White Plague Initial Trait Space Model

2019-2020 NSF EAGER Annual Report

Code for the initial trait space model and implementation.

Methods:

Our trait-space model applies an environmental filter (i.e. a white plague outbreak) to a reef community and then uses a hierarchical Bayesian model that incorporates intraspecific trait variation to predict the community assemblage after the environmental filter is applied. First, an ideal trait distribution for being resistant to the disease is created by taking the trait averages of the resistant species. Initially, only M. cavernosa values of Thalassobius mediterraneus abundance and Prophenyloxidase concentration were used to create the ideal trait distribution. This ideal, target trait distribution is essentially then environmental filter. Mclust, an R package for normal mixture modeling for model based clustering, classification, and density estimation, is then used to compute the probability of a trait given a species. Traits are drawn from the ideal traits distribution. This is the probability of a trait given the environment where the environment is the target traits. The posterior for the probability of a species given its traits and environment (target traits filter) is calculated. Finally, the posterior probability of a species given the ideal traits is determined by integrating out the traits. The model produces post-outbreak relative abundances of the species based on the resistant trait targets.

Code:

Packages needed

```
library(FD)
                    # gower dissimiliarity matrix and
#some other clustering stuff
library(ggplot2)
                    # plotting
library(vegan)
                    # stats
library(tidyverse) # data manipulation
library(cluster)
                 # clustering algorithms
library(factoextra) # clustering visualization
library(dendextend) # for making dendograms fancy
library(clValid)
# has the dunns index function for comparing cluster numbers
library(mclust)
# normal mixture modeling for model based clustering,
#classification, and density estimation
library(randomForest) # Random Forests functions
library(MASS)
```

Load in data

```
setwd("~/Dropbox/disease_trait_models/DiseaseTraitSpace_WP")
trait.data<-read.csv("Clustering_and_RF/traits_loggenes_env.csv",row.names=1)
#trait dataframes where each column is a different species
propheno<-read.csv("Trait_Space_Model/propheno.csv") #Prophenyloxidase concentration
thalmed<-read.csv("Trait_Space_Model/thalmed.csv") #Thalassobius mediterraneus</pre>
```

Step 1a.

Instead of an environmental gradient, we want an ideal trait distribution for being resistant. Initially using M. cavernosa traits as the ideal because it is the most resistant species

Get data for creating target traits distribution

```
#covariance
cov(propheno$Mcav.AntOx Prophenyloxidase,thalmed$Mcav.Bac thalassobius mediterraneus)
## [1] 0.0005296897
#propheno mean and variance
mean(propheno$Mcav.AntOx Prophenyloxidase)
## [1] 4.81985
var(propheno$Mcav.AntOx Prophenyloxidase)
## [1] 3.397597
#thalmed mean and variance
mean(thalmed$Mcav.Bac_thalassobius_mediterraneus)
## [1] 0.005633363
var(thalmed$Mcav.Bac_thalassobius_mediterraneus)
## [1] 5.477145e-06
Create the target traits distribution
reps < -c(1:10)
# defined early, but this is basically the number of simulations being run.
#propheno values first
covariance<-matrix(c(3.3976,0.0005,0.0005,0.00005),ncol=2)</pre>
means <-c(4.8199, 0.0056)
targets <-mvrnorm(n=1000,mu=means,Sigma=covariance)</pre>
#### Fit multivariate normal mixture model
pdf_targets <- mvn(modelName="XXX",targets,warn=TRUE)</pre>
#Computes the mean, covariance, and log-likelihood
#from fitting a single Gaussian to given data (univariate or multivariate normal).
#"XXX" for a general ellipsoidal Gaussian.
### Might want to change this eventually,
#try seeing what using Mclust instead looks like.
par_targets <- pdf_targets$parameters</pre>
```

Step 1b:

Use Mclust to compute the probability of a trait given the species

```
trait1<-trait.data$AntOx_Prophenyloxidase #set trait 1
trait2<-trait.data$Bac_thalassobius_mediterraneus #set trait 2
species<-trait.data$Species
Sp_data <- cbind(species,trait1,trait2)

cnat<-as.data.frame(Sp_data)%>%
    filter(species==1)
pdf_cnat<-Mclust(cnat[,2:3],warn=TRUE)
par_cnat<-pdf_cnat$parameters

mcav<-as.data.frame(Sp_data)%>%
    filter(species==2)
```

```
pdf_mcav<-Mclust(mcav[,2:3],warn=TRUE)</pre>
par_mcav<-pdf_mcav$parameters</pre>
oann<-as.data.frame(Sp_data)%>%
  filter(species==3)
pdf_oann<-Mclust(oann[,2:3],warn=TRUE)</pre>
par_oann<-pdf_oann$parameters
ofav<-as.data.frame(Sp data)%>%
  filter(species==4)
pdf_ofav<-Mclust(ofav[,2:3],warn=TRUE)</pre>
par_ofav<-pdf_ofav$parameters
past<-as.data.frame(Sp_data)%>%
  filter(species==5)
pdf_past<-Mclust(past[,2:3],warn=TRUE)</pre>
par_past<-pdf_past$parameters</pre>
ppor<-as.data.frame(Sp_data)%>%
  filter(species==6)
pdf_ppor<-Mclust(ppor[,2:3],warn=TRUE)</pre>
par_ppor<-pdf_ppor$parameters</pre>
ssid<-as.data.frame(Sp_data)%>%
  filter(species==7)
pdf ssid<-Mclust(ssid[,2:3],warn=TRUE)</pre>
par_ssid<-pdf_ssid$parameters
```

Step 2a:

Draw traits from the target traits distribution

```
#drawing samples from mixture densities fitted in Step 1A
N<- 100

trt_sample <- matrix(0,length(reps)*N,2)

trt_sample[1:N,] <- sim(pdf_targets$modelName,par_targets,N)[,2:3]

trt_sample[(N+1):(2*N),]<-sim(pdf_targets$modelName,par_targets,N)[,2:3]

trt_sample[(2*N+1):(3*N),]<-sim(pdf_targets$modelName,par_targets,N)[,2:3]

trt_sample[(3*N+1):(4*N),]<-sim(pdf_targets$modelName,par_targets,N)[,2:3]

trt_sample[(4*N+1):(5*N),]<-sim(pdf_targets$modelName,par_targets,N)[,2:3]

trt_sample[(5*N+1):(6*N),]<-sim(pdf_targets$modelName,par_targets,N)[,2:3]

trt_sample[(6*N+1):(7*N),]<-sim(pdf_targets$modelName,par_targets,N)[,2:3]

trt_sample[(8*N+1):(9*N),]<-sim(pdf_targets$modelName,par_targets,N)[,2:3]

trt_sample[(9*N+1):(10*N),]<-sim(pdf_targets$modelName,par_targets,N)[,2:3]</pre>
```

Probability of a trait given the "environment," where the environment is the target traits distribution

Step 2b.

Get the probability that a species has the sampled traits

```
#computing P(T/Sk) using Mclust done earlier
#note that in 2014 laughlin does the exp(trt_sample)
P_T_cnat<-dens(pdf_cnat$modelName,trt_sample,parameters=par_cnat)

P_T_oann<-dens(pdf_oann$modelName,trt_sample,parameters=par_oann)

P_T_ofav<-dens(pdf_ofav$modelName,trt_sample,parameters=par_ofav)

P_T_ppor<-dens(pdf_ppor$modelName,trt_sample,parameters=par_ppor)

P_T_past<-dens(pdf_past$modelName,trt_sample,parameters=par_past)

P_T_ssid<-dens(pdf_ssid$modelName,trt_sample,parameters=par_ssid)

P_T_mcav<-dens(pdf_mcav$modelName,trt_sample,parameters=par_mcav)

P_T_S <- cbind(P_T_cnat,P_T_mcav,P_T_oann,P_T_ofav,P_T_past,P_T_ppor,P_T_ssid)
summary(P_T_S)</pre>
```

```
##
      P T cnat
                 P_T_mcav
                                                    P_T_ofav
                                  P_T_oann
## Min. :0
             Min. :
                         0.00
                               Min. : 0.00000
                                                 Min.
                                                      :0
## 1st Qu.:0
             1st Qu.:
                         0.00
                               1st Qu.: 0.00000
                                                 1st Qu.:0
## Median :0
              Median :
                         0.00
                               Median : 0.00000
                                                 Median:0
## Mean
              Mean : 21.97
                               Mean
                                     : 0.05711
                                                 Mean
                                                        :0
         :0
## 3rd Qu.:0
              3rd Qu.:
                         0.00
                               3rd Qu.: 0.00000
                                                 3rd Qu.:0
## Max.
              Max. :7806.31
                                      :50.92245
         :0
                               Max.
                                                 Max.
##
      P_T_past
                        P_T_ppor
                                         P_T_ssid
## Min. : 0.00000
                    Min. :0.00000 Min.
                                            :0.000e+00
## 1st Qu.: 0.00000
                    1st Qu.:0.00000
                                      1st Qu.:0.000e+00
## Median : 0.00000
                     Median :0.00000
                                      Median :0.000e+00
## Mean
         : 0.08764
                            :0.00218
                                             :1.954e-33
                     Mean
                                      Mean
##
   3rd Qu.: 0.00000
                     3rd Qu.:0.00000
                                      3rd Qu.:0.000e+00
## Max.
          :15.15690
                     Max.
                            :1.26374
                                      Max.
                                             :1.385e-30
```

Multiply likelihood by a prior to get the posterior

```
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 0.0000 0.0000 0.0000 0.7962 0.0000 273.2209
```

Step 2C.

Get the Posterior for the probability of a species given traits and environment

```
#computing P(Sk/T,E) using Bayes theorem

P_S_T_E = matrix(0,dim(P_T_S)[1],7)

for (i in 1:dim(P_T_S)[1]){
    P_S_T_E[i,] = exp(log(P_T_S_pr[i,]) - log(P_T_S_pr_sum[i]))
    } #using log
P_S_T_E[is.nan(P_S_T_E)] <- 0</pre>
```

Step 2D

Get the posterior probability of a species given the target resistant traits, integrate out the traits.

```
#P(Sk/T) by integrating out T's
P_S_E_all = matrix(0,length(reps)*N,7)
P_S_E_unnorm = matrix(0,length(reps),7) #unnormalised P_S_E
P_S_E = matrix(0,length(reps),7)

#before MC integration (with log)
for (i in 1:dim(P_S_E_all)[1]){
    P_S_E_all[i,]=exp(log(P_T_E[i])+log(P_S_T_E[i,]))}
#MC integration and normalisation
c=1
for (k in 1:length(reps)){
    c=(k-1)*N+1
    P_S_E_unnorm[k,]=apply(P_S_E_all[c:(c+(N-1)),],2,mean) #MC
    P_S_E[k,]=P_S_E_unnorm[k,]/sum(P_S_E_unnorm[k,]) #normalisation
}
apply(P_S_E,1,sum) ### check that probs sum to one
```

```
## [1] 1 1 1 1 1 1 1 1 1 1 1 P_S_E
```

```
[,1]
##
                   [,2]
                                 [,3] [,4]
                                                [,5]
                                                             [,6]
                                                                        [,7]
##
  [1,]
            0 0.6292513 4.212077e-03
                                         0 0.1656253 1.049390e-02 0.19041742
## [2,]
            0 0.6962527 1.054080e-214
                                        0 0.1767030 5.296881e-03 0.12174748
## [3,]
           0 0.6769211 4.674194e-04
                                        0 0.1422834 2.391717e-38 0.18032810
## [4,]
           0 0.6978246 8.550812e-13
                                        0 0.1574034 7.713791e-03 0.13705821
           0 0.7052006 2.456588e-03
## [5,]
                                        0 0.1116279 4.857254e-03 0.17585765
## [6,]
           0 0.7545862 2.167446e-03
                                        0 0.1136071 9.636522e-03 0.12000266
```

```
[7,]
            0 0.7088347 1.508488e-03
                                         0 0.1830892 9.594835e-03 0.09697278
##
                                         0 0.1520579 1.187641e-02 0.11206783
##
   [8,]
            0 0.7208854 3.112494e-03
   [9,]
                                         0 0.1454341 7.449528e-03 0.18534772
##
            0 0.6559715 5.797136e-03
## [10,]
            0 0.6841227
                         1.135931e-03
                                         0 0.1600854 1.310224e-02 0.14155373
### probs
```

Post-Outbreak abundances based on 'resistant trait targets'

Sp. abundances post-outbreak, mcav is target

