Produce ternary plots of elasticities

2015-10-08

Here we produce a ternary plot a la Silvertown & Franco (1993) with various life history traits such as mean life expectancy, population growth rate or reactivity as the "fourth" dimension. We will use Caswell (2001) formulation of mean life expectancy from the fundamental matrix (N), and the packages popdemo for the reactivity:

```
require(popdemo)
```

> Loading required package: popdemo

This is the function to calculate mean life expectancy from Caswell (2001):

```
meanLifeExpectancy <- function(matU = matU, startLife = 1){
    uDim=dim(matU)[1]
    N = solve(diag(uDim[startLife])-matU)
    eta = colSums(N)[startLife]
    return(eta)
}</pre>
```

As an example for the chosen data, subset COMADRE to studies with a matrix dimension >= 3, that represent mean, unmanipulated conditions duration > 3 years, where sexual reproduction has been modeled explicitly, the matrices are split into U, F and C, and there are no issues with stage-specific survival >1.

Use the row names from the subsetted dataframe to subset the matrices.

```
keep <- as.numeric(rownames(tempMetadata))</pre>
```

Define the object tempMat as the list object containing matrices in the same order that their metadata appears in tempMetadata.

```
tempMat <- comadre$mat[keep]</pre>
```

These matrices can now be analyzed by applying functions in a loop, or by using lapply.

To calculate elasticities, population growth rate, reactivity and mean life expectancy for the subset matrices, first create an empty data.frame to accommodate the output:

Use the following function to calculate element-level perturbations:

```
matrixElementPerturbation <- function(matU, matF, matC=NULL,pert=0.001){
 matA=matU+matF+matC
  aDim=dim(matA)[1]
 fakeA=matA
  sensA=elasA=matrix(NA,aDim,aDim)
 lambda=Re(eigen(matA)$values[1])
 propU=matU/matA
   propU[is.nan(propU)]=NA
   propProg=propRetrog=propU
   propProg[upper.tri(propU,diag=T)]=NA
   propRetrog[lower.tri(propU,diag=T)]=NA
   propStasis=matrix(diag(aDim)*diag(propU),aDim,aDim)
 propF=matF/matA
    propF[is.nan(propF)]=NA
  propC=matC/matA
   propC[is.nan(propC)]=NA
 for (i in 1:aDim){
    for (j in 1:aDim){
       fakeA=matA
       fakeA[i,j]=fakeA[i,j]+pert
       lambdaPert=eigen(fakeA)$values[1]
       sensA[i,j]=(lambda-lambdaPert)/(matA[i,j]-fakeA[i,j])
   }
 }
  sensA=Re(sensA)
  elasA=sensA*matA/lambda
  out = data.frame("SStasis"=NA, "SProgression"=NA, "SRetrogression"=NA,
                   "SFecundity"=NA, "SClonality"=NA, "EStasis"=NA, "EProgression"=NA,
                   "ERetrogression"=NA, "EFecundity"=NA, "EClonality"=NA)
    out$SStasis=sum(sensA*propStasis,na.rm=T)
    out$SRetrogression=sum(sensA*propRetrog,na.rm=T)
    out$SProgression=sum(sensA*propProg,na.rm=T)
    out$SFecundity=sum(sensA*propF,na.rm=T)
    out$SClonality=sum(sensA*propC,na.rm=T)
    out$EStasis=sum(elasA*propStasis,na.rm=T)
    out$EProgression=sum(elasA*propProg,na.rm=T)
    out$ERetrogression=sum(elasA*propRetrog,na.rm=T)
    out$EFecundity=sum(elasA*propF,na.rm=T)
    out$EClonality=sum(elasA*propC,na.rm=T)
 return(out)
}
```

Now we can use a loop to examine each matrix:

```
for (i in 1:length(tempMat)){
  tryCatch({
    matA=tempMat[[i]]$matA
    matU=tempMat[[i]]$matU
    matF=tempMat[[i]]$matF
  matC=tempMat[[i]]$matC
```

Group elasticities of population growth rate to various demographic processes into three main axes:

```
output$S=output$EStasis+output$ERetrogression
output$G=output$EProgression
output$R=output$EFecundity+output$EClonality
```

Scale to 1 the coordinates of each point - this is necessary due to possible rounding issues, although note that the function to create the ternary plot below can do this automatically with the argument scale.

```
output$S=output$S/rowSums(output[,c("S","G","R")])
output$G=output$G/rowSums(output[,c("S","G","R")])
output$R=output$R/rowSums(output[,c("S","G","R")])
```

Eliminate the couple of MPMs where the code did not run correctly, as it produced all NAs:

```
output=output[-which(is.na(output$eta)),]
```

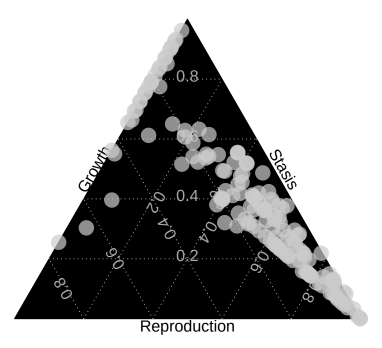
Plot the locations of the chosen matrices in a preliminary ternary plot, for which the following libraries will also be necessary

```
library(vcd)
```

> Loading required package: grid

```
library(scales)
```

Preliminary plot



Color-code the points in the ternary plot according to range of lambda, eta and reactivity, respectively, on different plots:

```
lambdaData <- output[which(log(output$lambdas)<=2),]</pre>
x_norm <- log(lambdaData$lambda)</pre>
minL<-min(lambdaData$lambda)</pre>
maxL<-max(lambdaData$lambda)</pre>
x_norm <- (lambdaData$lambda-minL)/(maxL-minL)</pre>
col_fun <- colorRamp(c("white","yellow","orange","red","dark red"))</pre>
rgb_cols <- col_fun(x_norm)</pre>
colsLambda <- rgb(rgb_cols, maxColorValue = 256)</pre>
etaData<-output
etaData$etalog<-log(etaData$eta)</pre>
minE<-min(etaData$etalog)</pre>
maxE<-max(etaData$etalog)</pre>
x_norm <- (etaData$etalog-minE)/(maxE-minE)</pre>
rgb_cols <- col_fun(x_norm)</pre>
colsEta <- rgb(rgb_cols, maxColorValue = 256)</pre>
reactData<-output[which(log(output$react)<=5),]</pre>
reactData$reactlog<-log(reactData$react)</pre>
minR <- min(output$react)</pre>
maxR <- max(output$react)</pre>
x_norm <- (output$react-minR)/(maxR-minR)</pre>
rgb_cols <- col_fun(x_norm)</pre>
colsReact <- rgb(rgb_cols, maxColorValue = 256)</pre>
```

And next plot all three ternary plots. The legend will need the library fields

```
library(fields)
```

```
> Loading required package: spam
```

```
> Spam version 1.0-1 (2014-09-09) is loaded.
> Type 'help( Spam)' or 'demo( spam)' for a short introduction
> and overview of this package.
> Help for individual functions is also obtained by adding the
> suffix '.spam' to the function name, e.g. 'help( chol.spam)'.
>
> Attaching package: 'spam'
>
> The following objects are masked from 'package:base':
>
> backsolve, forwardsolve
>
> Loading required package: maps

We can use the same colour palette for each of the plots:

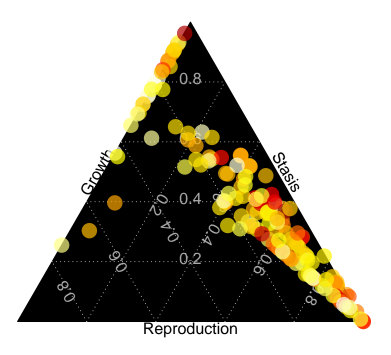
colCode <- colorRampPalette(c("white","yellow","orange","red","dark red"))(n = 999)

zr <- range(c(lambdaData$lambda, na.rm = TRUE))
ternaryplot(lambdaData[,c("R","S","G")],scale=1,col=alpha(colsEta,0.7),bg="black",")</pre>
```

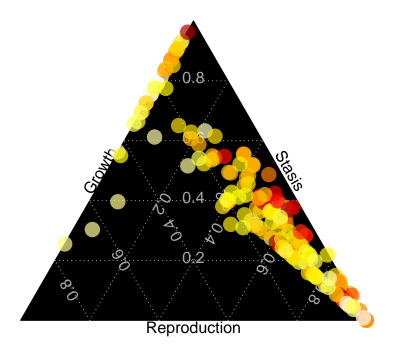
dimnames=c("Stasis", "Growth", "Reproduction"), dimnames_position="edge",

main=expression(paste("Population growth rate - ", lambda)))

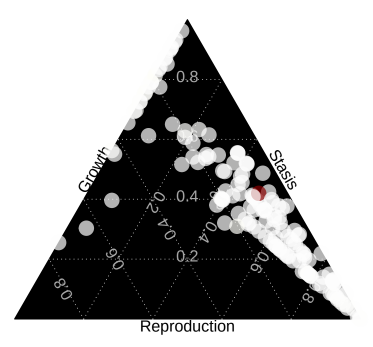
Population growth rate $-\lambda$



Mean life expectancy – η_e



Reactivity – $||\hat{A}||_1$



One might add the color scale legend using the following code to add to the same plot:

```
## Warning in par(new = TRUE, pty = "m", plt = smallplot, err = -1): calling ## par(new=TRUE) with no plot
```

