Working notes ®

ggplot2 implementation of the graphical functions of the ade4 package (in working!)

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1 Motivations

The objectives of this document are to propose elements and alternatives for ggplot2 implementation ([5]) of the graphical function from R package ADE-4 ([2, 3, 4]). why ggplot2?

2 Gestion of the limits

A function getLimits is written to extract the limits of the axes x and y as done in the R package ade4.

```
getLimits <- function (dfxy, xax=1, yax=2,include.origin=TRUE,origin=c(0,0)){</pre>
            df <- data.frame(dfxy)</pre>
            if ((xax < 1) || (xax > ncol(df)))
stop("Non convenient selection for xax")
            if ((yax < 1) \mid | (yax > ncol(df)))
                      stop("Non convenient selection for yax")
           x <- df[, xax]
y <- df[, yax]
            x1 <- x
if (include.origin)
            x1 <- c(x1, origin[1])
x1 <- c(x1 - diff(range(x1)/10), x1 + diff(range(x1))/10)
            xlim <- range(x1)</pre>
            у1 <- у
            if (include.origin)
            y1 <- c(y1, origin[2])
y1 <- c(y1 - diff(range(y1)/10), y1 + diff(range(y1))/10)
            ylim <- range(y1)</pre>
            return(list(xlim=xlim, ylim=ylim))
 }
Example for a futur prototype of the function ggade (or ggscatter?)
  require(ggplot2)
  ggade <- function(dfxy,xax=1,yax=2,...,include.origin=TRUE,origin=c(0,0)){
  yxlim <- getLimits(dfxy,xax=xax,yax=yax,include.origin=include.origin,origin=origin)
  ggplot(....) + coord_cartesian(xlim=yxlim$xlim,ylim=yxlim$ylim) + coord_fixed(ratio=1)</pre>
```

3 Representations of the variables

```
data(deug)
deug0 <- dudi.pca(deug$tab, center = deug$cent, scale = FALSE, scan = FALSE)
deug1 <- dudi.pca(deug$tab, center = TRUE, scale = TRUE, scan = FALSE)</pre>
```

```
require(ggplot2)
gg <- ggplot(data=data.frame(eig=deug1$eig,nf=1:length(deug1$eig)), aes(x=nf, y=eig))
gg <- gg + geom_bar(stat="identity") + ggtitle("Eigenvalues from PCA")
gg <- gg + ylab("Eigenvalues") + xlab("axis") + theme_light()
gg</pre>
```

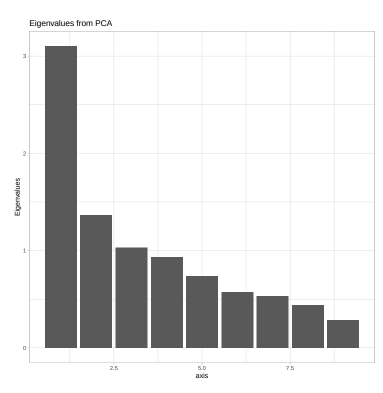


Figure 1: Representation of the Eigenvalues from PCA on correlation matrix

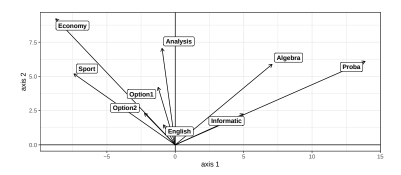


Figure 2: Representation of the variables with correlation circle for PCA on covariance

```
require(cowplot)
require(ggrepe1)
require(ggrepe1)
require(ggforce)
auxi <- deug1$co
auxi$label <- rownames(auxi)
ggx <- ggplot(data=auxi,aes(Comp1,Comp2,label=label))
ggx <- ggx + geom_hline(yintercept = 0)+geom_vline(xintercept = 0)
ggx <- ggx + xlab("axis 1") + ylab("axis 2")
ggx <- ggx + geom_circle(data=data.frame(x0=0,y0=0),aes(x0=x0, y0=y0, r=1),inherit.aes = FALSE)
ggx <- ggx + geom_segment(aes(x=0,xend =Comp1, y=0,yend = Comp2),arrow = arrow(length = unit(0.2,"cm")))
ggx <- ggx + geom_label_repel(size = 3.5,segment.alpha=0.7,segment.color = "darkgrey",fontface = 'bold',col="
ggx <- ggx + theme_bw() + coord_fixed(ratio=1)
ggx</pre>
```

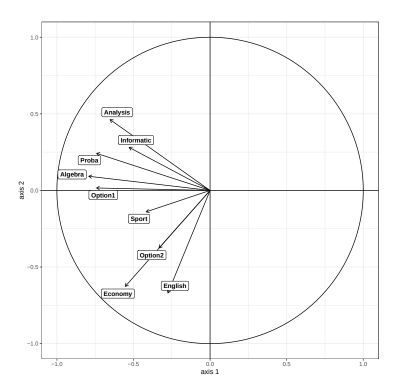


Figure 3: Representation of the variables with correlation circle for PCA on correlation

4 Representations of samples

representation of the samples on the first factorial plan

```
require(ggplot2)
require(ggrepel)
auxi <- deug1$1i
auxi$label <- rownames(auxi)
gg <- ggplot(auxi,aes(Axis1,Axis2,label=label))
gg <- gg + geom_hline(yintercept = 0) + geom_vline(xintercept = 0)
gg <- gg + geom_point(shape=19,size=3)
gg <- gg + geom_label_repel(size = 3,segment.alpha=0.7,segment.color = "darkgrey")
gg <- gg + theme_bw()+ coord_fixed(ratio=1)
gg</pre>
```

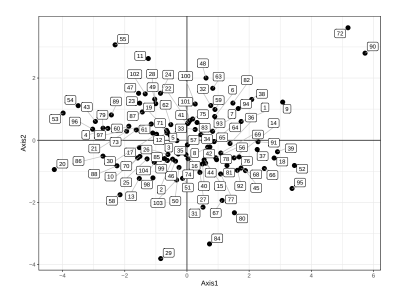


Figure 4: Representation of the observation of the first vectorial plan

5 Representations of samples by group

5.1 separated Representations of samples by group

Representation of the samples on the first factorial plan

```
require(ggplot2)
require(ggrepel)
auxi <- deug1$li
auxi$label <- rownames(auxi)
auxi$group <- substring(as.character(deug$result),1,1)
gg <- ggplot(auxi,aes(Axis1,Axis2,label=label))
gg <- gg + geom_hline(yintercept = 0) + geom_vline(xintercept = 0)
gg <- gg + geom_point(shape=19,size=3)
gg <- gg + geom_label_repel(size = 3,segment.alpha=0.7,segment.color = "darkgrey")
gg <- gg + theme_bw() + coord_fixed(ratio=1) + facet_wrap(~ group)
gg</pre>
```

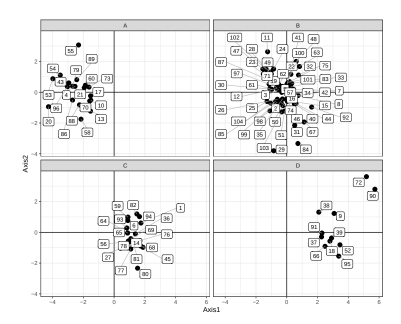


Figure 5: Representation of the observation of the first vectorial plan stratified by group.

```
require(ggplot2)
require(ggrepel)
auxi <- deug1$li
auxi$label <- rownames(auxi)
auxi$group <- substring(as.character(deug$result),1,1)
gg <- ggplot(auxi,aes(Axis1,Axis2,label=label,color=group))
gg <- gg + geom_hline(yintercept = 0) + geom_vline(xintercept = 0)
gg <- gg + geom_point(shape=19,size=3)
gg <- gg + geom_label_repel(size = 3,segment.alpha=0.7,segment.color = "darkgrey")
gg <- gg + theme_bw() + coord_fixed(ratio=1) + facet_wrap(~ group)
gg</pre>
```

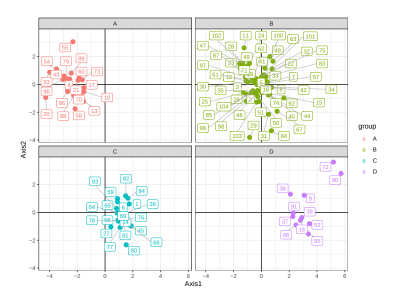


Figure 6: Representation of the observation of the first vectorial plan stratified by group (with color).

5.2 Ellipses

```
require(ggplot2)
require(ggrepel)
auxi <- deug1$li
auxi$label <- rownames(auxi)
auxi$group <- substring(as.character(deug$result),1,1)
gg <- ggplot(auxi,aes(Axis1,Axis2,label=label,color=group))
gg <- gg + geom_hline(yintercept = 0) + geom_vline(xintercept = 0)
gg <- gg + geom_point(shape=19,size=3)
gg <- gg + stat_ellipse(type = "norm",level=0.66)
gg <- gg + geom_label_repel(size = 3,segment.alpha=0.7,segment.color = "darkgrey")
gg <- gg + theme_bw() + coord_fixed(ratio=1)
gg</pre>
```

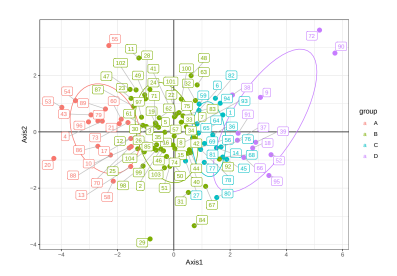


Figure 7: Representation of the observation of the first vectorial plan with ellpise of Inertia for each group

```
require(ggplot2)
require(ggrepel)
auxi <- deug1$li
auxi$label <- rownames(auxi)
auxi$group <- substring(as.character(deug$result),1,1)
gg <- ggplot(auxi,aes(Axis1,Axis2,label=label,color=group))
gg <- gg + geom_hline(yintercept = 0) + geom_vline(xintercept = 0)
gg <- gg + geom_point(shape=19,size=3)
gg <- gg + stat_ellipse(type = "norm",level=0.66)
gg <- gg + geom_label_repel(size = 3,segment.alpha=0.7,segment.color = "darkgrey")
gg <- gg + theme_bw() + coord_fixed(ratio=1) + facet_wrap(~group)</pre>
```

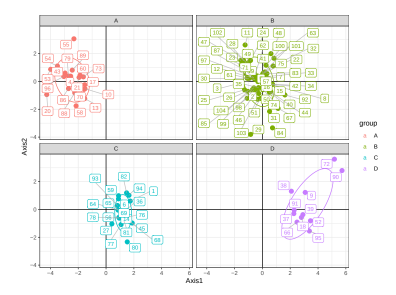


Figure 8: Representation of the observation of the first vectorial plan with ellpise of Inertia for each group

5.3 Ellipses and stars

For this representation, we need to compute the position of the gravity center of each ellipse. This operation was performed by the R function prep.tab.class. The implementation of this function is given below:

```
prep.tab.class <- function(x,fac,varnames=c("Axis1","Axis2"),rm.X=TRUE,...){
    x <- as.data.frame(x)
    x$fac <- factor(x[,fac])
    w <- tab.class(x[,varnames],fac=x$fac)
    colnames(w) <- c("Axis1","Axis2")
    if(rm.X){
        w$label <- gsub("X","",rownames(w)) #levels(x$fac)
}else{
        w$label <- rownames(w)
}
rownames(w) <- as.character(w$label)</pre>
```

```
x$cAxis1 <- w$Axis1[match(as.character(x$fac),w$label)]
x$cAxis2 <- w$Axis2[match(as.character(x$fac),w$label)]
return(list(data=x,center=w))

require(ggrepel)
auxi <- deugi$li
auxi$label <- rownames(auxi)
auxi$group <- substring(as.character(deug$result),1,1)
w <- prep.tab.class(auxi,varnames=c("Axis1","Axis2"),fac="group",rm.X=FALSE)
datax <- w$data
centerx <- w$center
gg <- ggplot(datax,aes(x=Axis1,y=Axis2,col=group,group=group))
gg <- gg + geom_hline(yintercept = 0) + geom_vline(xintercept = 0)
gg <- gg + stat_ellipse(type = "norm",level=0.66,lwd=1,show.legend = FALSE)
gg <- gg + geom_point(data=catax,aes(x=Axis1,y=Axis2,col=label),inherit.aes = FALSE,size=0)
gg <- gg + geom_point(data=centerx,aes(x=Axis1,y=Axis2,col=label),inherit.aes = FALSE,size=0)
gg <- gg + geom_point(shape=19,size=2,show.legend = FALSE)
# gg <- gg + geom_label_repel(data=centerx,aes(x=Axis1,y=Axis2,col=label,label=label),size = 3,
# segment.alpha=0.7,segment.color = "darkgrey",fontface = 'bold',inherit.aes = FA
g <- gg + geom_label(data=centerx,aes(x=Axis1,y=Axis2,col=label,label=label),size = 3,
fontface = 'bold',inherit.aes = FALSE)
gg <- gg + theme_light() +theme(legend.position = "none") + coord_fixed(ratio=1)
gg <- gg + theme_light() +theme(legend.position = "none") + coord_fixed(ratio=1)</pre>
```

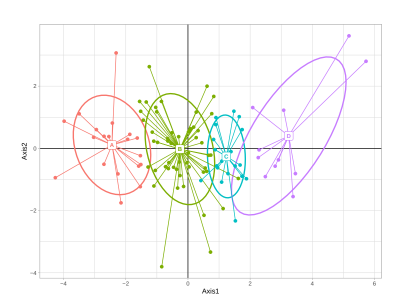


Figure 9: Representation of the observation of the first vectorial plan with ellpise of Inertia for each group

5.4 Chull representation

The following code is based on the vignette related to the extending ggplot2 functions (https://cran.r-project.org/web/packages/ggplot2/vignettes/extending-ggplot2.htm). The additional functions for the computation necessary to the Convex hull representation are given below:

The representation for one group is given below:

```
require(ggplot2)
# data
auxi <- deug1$li
auxi$label <- rownames(auxi)
auxii$group <- substring(as.character(deug$result),1,1)
# Find the convex hull of the points being plotted
# Define the scatterplot
gg <- ggplot(auxi,aes(Axis1,Axis2,col=group))
gg <- gg + geom_hline(yintercept = 0) + geom_vline(xintercept = 0)
gg <- gg + geom_point(shape=19,size=3)
gg <- gg + stat_chull(fill=NA)
gg <- gg + theme_bw() + coord_fixed(ratio=1)
gg</pre>
```

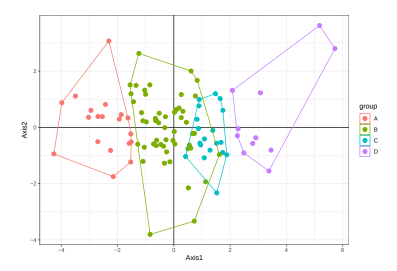


Figure 10: Chull representation of the observation of the first vectorial plan.

```
require(ggplot2)
# data
auxi <- deug1$li
auxi$label <- rownames(auxi)
auxii$group <- substring(as.character(deug$result),1,1)
# Find the convex hull of the points being plotted
# Define the scatterplot
gg <- ggplot(auxi,aes(Axis1,Axis2,col=group,fill=group))
gg <- gg + geom_hline(yintercept = 0) + geom_vline(xintercept = 0)
gg <- gg + geom_point(shape=19,size=3)
gg <- gg + stat_chull(alpha=0.5)
gg <- gg + theme_bw() + coord_fixed(ratio=1)
gg</pre>
```

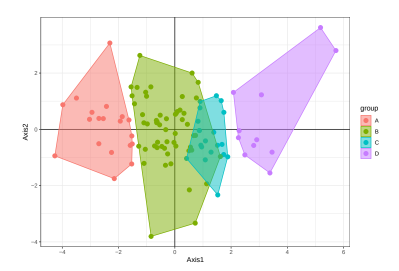


Figure 11: Chull representation of the observation of the first vectorial plan.

6 2D density

```
require(ggplot2)
require(ggrepel)
auxi <- deug1$li
auxi$label <- rownames(auxi)
auxi$group <- deug$result
gg <- ggplot(auxi,aes(Axis1,Axis2,label=label))
gg <- gg + geom_hline(yintercept = 0) + geom_vline(xintercept = 0)
gg <- gg + geom_point(shape=19,size=3)
gg <- gg + stat_density2d(col="blue")
gg <- gg + theme_bw() + coord_fixed(ratio=1)
gg</pre>
```

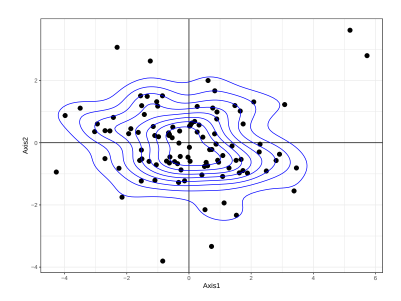


Figure 12: Representation of the observation with 2-D kernel density

```
require(ggplot2)
require(ggrepe1)
auxi <- deugi$li
auxi$label <- rownames(auxi)
auxi$group <- substring(as.character(deug$result),1,1)
gg <- ggplot(auxi,aes(Axis1,Axis2,label=label,col=group))
gg <- gg + geom_hline(yintercept = 0) + geom_vline(xintercept = 0)
gg <- gg + geom_point(shape=19,size=3)
gg <- gg + stat_density2d(col="blue")
gg <- gg + theme_bw() + coord_fixed(ratio=1)
gg</pre>
```

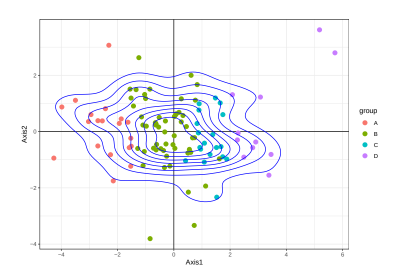


Figure 13: Representation of the observation with 2-D kernel density

```
require(ggplot2)
require(ggrepel)
auxi <- deug1$1i
auxi$label <- rownames(auxi)
auxi$group <- substring(as.character(deug$result),1,1)
gg <- ggplot(auxi,aes(Axis1,Axis2,label=label,col=group))
gg <- gg + geom_hline(yintercept = 0) + geom_vline(xintercept = 0)
gg <- gg + geom_point(shape=19,size=3)
gg <- gg + stat_density2d(col="blue")
gg <- gg + theme_bw() + coord_fixed(ratio=1) + facet_wrap(~group)
gg</pre>
```

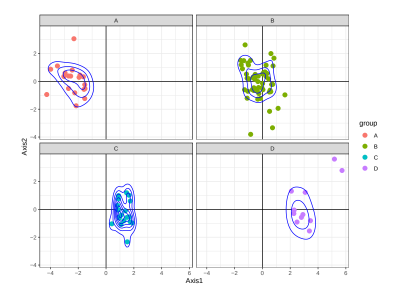


Figure 14: Representation of the observation with 2-D kernel density

7 Adding a picture

The functions from the R package imager ([1]) can be used To add picture on ggplot graphics (https://cran.r-project.org/web/packages/imager/vignettes/gettingstarted.html).

```
library(ggplot2)
library(dplyr)
library(imager)
parrots <- load.image("/export/scratch/R/library/imager/extdata/parrots.png")
df <- as.data.frame(parrots,wide="c") %>% mutate(rgb.val=rgb(c.1,c.2,c.3))
df$y <- rev(df$y)
gg <- ggplot(df,aes(x,y)) + geom_raster(aes(fill=rgb.val)) + scale_fill_identity()
gg <- gg + xlab("x") + ylab("y")
gg <- gg + theme_bw() + coord_fixed(ratio=1)
gg</pre>
```

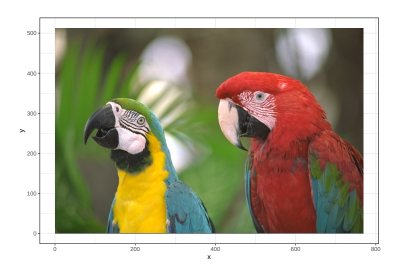


Figure 15: Representation of "parrots" picture from R package imager.

Test for PNM format with imager ... resultats boff (?!)

```
library(ggplot2)
library(dplyr)
library(imager)
#photo1 <- load.image("/export/scratch/R/library/ade4/inst/pictures/buterfly.pnm")
maps1 <- load.image("/export/scratch/R/library/ade4/pictures/butterfly.pnm")
df <- as.data.frame(maps1)
# black=#000000 and white = #FFFFFF
df$rgb.val <- ifelse(df$value==0,'#000000','#FFFFFF')
df$y <- rev(df$y)
gg <- ggplot(df,aes(x,y)) + geom_raster(aes(fill=rgb.val)) + scale_fill_identity()
gg <- gg + xlab("x") + ylab("y")
gg <- gg + theme_bw() + coord_fixed(ratio=1)
gg</pre>
```

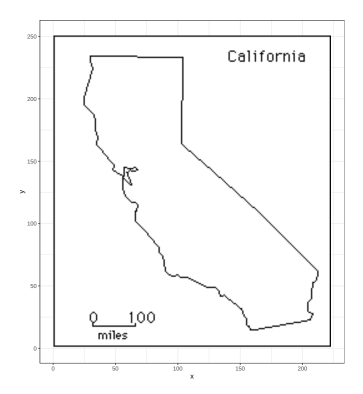


Figure 16: Representation of the map associated with butterfly data from R package ade4.

For the color code see https://www.nceas.ucsb.edu/sites/default/files/2020-04/colorPaletteCheatsheet.pdf. The previous results is not very good. we try to used the R package magick https://docs.ropensci.org/magick/articles/intro.html). The code is given below:

```
require(magick)
maps2 <- image_read("/export/scratch/R/library/ade4/pictures/butterfly.pnm")</pre>
```

add points and information

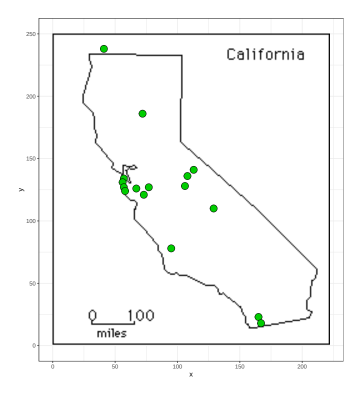


Figure 17: Representation of the map associated with butterfly data from R package ade4.

8 Complex figures and Examples

The complex figures are aggregated with the package cowplot ([6]).

8.1 Heatmap and PCA

preliminary analysis

```
tabenv <- butterfly$envir
tabgen <- butterfly$genet
pcaenv <- dudi.pca(tabenv,nf=2,scannf=FALSE)
pcagen <- dudi.pca(tabgen,nf=2,scannf=FALSE)</pre>
```

preparation of the graphics related to the PCAs of genetic features and environmental variables.

```
require(ggplot2)
  require(ggrepel)
  require(ggforce)
### variable representation
  auxi <- pcaenv$co
  auxi$label <- rownames(auxi)
ggx <- ggplot(data=auxi,aes(Comp1,Comp2,label=label))</pre>
  ggx <- ggx + geom_hline(yintercept = 0)+geom_vline(xintercept = 0)
ggx <- ggx + xlab("axis 1") + ylab("axis 2")
ggx <- ggx + geom_circle(data=data.frame(x0=0,y0=0),aes(x0=x0, y0=y0, r=1),inherit.aes = FALSE)</pre>
  ggx <- ggx + geom_label_repel(size = 3.5,segment.alpha=0.7,segment.color = "darkgrey",</pre>
  ggVarEnv <- ggx + theme_bw() + coord_fixed(ratio=1)
  auxi <- pcagen$co
  ggx <- ggx + geom_hline(yintercept = 0)+ggx <- ggx + geom_hline(yintercept = 0)+ggx <- ggx + geom_hline(yintercept = 0)+geom_vline(xintercept = 0)</pre>
  auxi <- pcaenv$li
  auxi$label <- rownames(auxi)
  gg <- ggplot(auxi,aes(Axis1,Axis2,label=label))</pre>
  gg <- ggpriot(aux1,aes(ax1s1,ax1s2,label=label))
gg <- gg + geom_hline(yintercept = 0) + geom_vline(xintercept = 0)
gg <- gg + geom_point(shape=19,size=3)
gg <- gg + geom_label_repel(size = 3,segment.alpha=0.7,segment.color = "darkgrey")
ggSampEnv <- gg + theme_bw()+ coord_fixed(ratio=1)</pre>
  auxi <- pcagen$li
  auxi$label <- rownames(auxi)</pre>
  gg <- ggplot(auxi,aes(Axis1,Axis2,label=label))</pre>
  gg <- gg + geom_hline(yintercept = 0) + geom_vline(xintercept = 0)
  gg <- gg + geom_point(shape=19,size=3)
  gg <- gg + geom_label_repel(size = 3,segment.alpha=0.7,segment.color = "darkgrey")
ggSampGen <- gg + theme_bw()+ coord_fixed(ratio=1)
## eigenvalues
  gg <- ggplot(data=data.frame(eig=pcaenv$eig,nf=1:length(pcaenv$eig)), aes(x=nf, y=eig))
gg <- gg + geom_bar(stat="identity") + ggtitle("Eigenvalues from PCA")
ggEigEnv <- gg + ylab("Eigenvalues") + xlab("axis") + theme_light()</pre>
  gg <- ggplot(data=data.frame(eig=pcagen$eig,nf=1:length(pcagen$eig)),
gg <- gg + geom_bar(stat="identity") + ggtitle("Eigenvalues from PCA")
ggEigGen <- gg + ylab("Eigenvalues") + xlab("axis") + theme_light()</pre>
                                                                                               aes(x=nf, y=eig))
preparation of the heatmap representation
  require(ComplexHeatmap)
  colgradient <- colorRamps::matlab.like2(25)</pre>
  top_annotation =HeatmapAnnotation(df=as.data.frame(tabenv)))
```

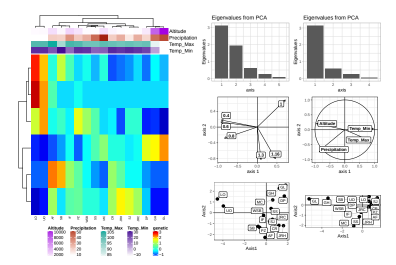


Figure 18: Representation of the preliminary analysis of butterfly data from R package ade4.

8.2 Coinertia analysis

8.3 K-table representations

9 Conclusion

References

- [1] Simon Barthelme. imager: Image Processing Library Based on 'CImg', 2019. R package version 0.41.2.
- [2] D. Chessel, A.B. Dufour, and J. Thioulouse. The ade4 package-I- One-table methods. $R\ News,\ 4:5-10,\ 2004.$
- [3] S. Dray and A.B. Dufour. The ade4 package: implementing the duality diagram for ecologists. *Journal of Statistical Software*, 22(4):1–20, 2007.
- [4] S. Dray, A.B. Dufour, and D. Chessel. The ade4 package-II: Two-table and K-table methods. R News, 7(2):47-52, 2007.
- [5] Hadley Wickham. ggplot2: Elegant Graphics for Data Analysis. Springer-Verlag New York, 2016.
- [6] Claus O. Wilke. cowplot: Streamlined Plot Theme and Plot Annotations for 'ggplot2', 2019. R package version 0.9.4.

10 Appendix

```
print(sessionInfo(),locale=FALSE)
R version 3.6.3 (2020-02-29)
Platform: x86_64-pc-linux-gnu (64-bit)
Running under: Linux Mint 18.3
Matrix products: default
BLAS: /usr/lib/openblas-base/libblas.so.3
LAPACK: /usr/lib/libopenblasp-r0.2.18.so
Random number generation:
RNG: Mersenne-Twister
Normal: Inversion
Sample: Rounding
attached base packages:
                              parallel stats base
 [1] grid datasets
[9] grDevices methods
                                                        graphics utils
                 datasets
                                                                                 stats4
                                                                                              tools
other attached packages:
 [1] ComplexHeatmap_2.0.0 magick_2.0
                                                           imager_0.41.2
 [4] magrittr_1.5
                                dplyr_1.0.2
                                                           ggforce_0.2.2
 [7] ggrepel_0.8.1
                                cowplot_0.9.4
                                                           ggplot2_3.3.2
[10] knitr_1.23
                                pixmap_0.4-11
                                                           ade4_1.7-15
[13] RColorBrewer_1.1-2
[16] GenomeInfoDb_1.20.0
[19] BiocGenerics_0.30.0
                                rtracklayer_1.44.0
                                                           GenomicRanges_1.36.0
                                IRanges_2.18.0
                                                           S4Vectors_0.22.1
loaded via a namespace (and not attached):
 [1] Rcpp_1.0.5
                                         circlize_0.4.8
 [3] lattice_0.20-38
                                         png_0.1-7
 [5] Rsamtools_2.0.0
                                         Biostrings_2.52.0
 [7] digest_0.6.22
                                         R6_2.4.0
 [9] tiff_0.1-5
                                         plyr_1.8.4
[11] pillar_1.4.7
                                         GlobalOptions_0.1.1
[13] zlibbioc_1.30.0
                                         rlang_0.4.9
[15] rstudioapi_0.10
                                         GetoptLong_0.1.7
[17] Matrix_1.2-17
                                         bmp_0.3
[19] labeling_0.3
                                         BiocParallel_1.18.1
[21] stringr_1.4.0
[23] RCurl_1.95-4.12
                                         igraph_1.2.4.1
                                         polyclip_1.10-0
                                         DelayedArray_0.10.0
[25] munsell_0.5.0
[27] compiler_3.6.3
                                         xfun_0.10
[29] pkgconfig_2.0.3
                                         shape_1.4.4
[31] readbitmap_0.1.5
                                         tidyselect_1.1.0
[33] SummarizedExperiment_1.14.0 tibble_3.0.4
[35] GenomeInfoDbData_1.2.1 matrixStats_(
[37] XML_3.98-1.20 crayon_1.3.4
                                         matrixStats_0.55.0
[39] withr_2.1.2
[41] MASS_7.3-51.4
                                         GenomicAlignments_1.20.0
                                         bitops_1.0-6
                                         lifecycle_0.2.0
[43] gtable_0.3.0
[45] scales_1.0.0
                                         stringi_1.4.3
[47] farver_1.1.0
[49] colorRamps_2.3
                                         XVector_0.24.0 ellipsis_0.3.0
[51] generics_0.0.2
                                         vctrs_0.3.5
[53] rjson_0.2.20
                                         Biobase_2.44.0
[55] glue_1.4.2
                                         tweenr_1.0.1
[57] purrr_0.3.4
[59] clue_0.3-57
                                         jpeg_0.1-8.1
                                         colorspace_1.4-1
[61] cluster_2.1.0
                                         isoband_0.2.3
  options(encoding="latin1",prompt="> ", continue=" ", width = 85)
> #save.image()
```