Package ‘aPCoA’

December 13, 2021

Type Package
Title Covariate Adjusted PCoA Plot
Version 1.3
Date 2021-12-12
Author Yushu Shi
Maintainer Yushu Shi <shiyushu2006@gmail.com>
Description In fields such as ecology, microbiology, and genomics, non-Euclidean distances are widely applied to describe pairwise dissimilarity between samples. Given these pairwise distances, principal coordinates analysis (PCoA) is commonly used to construct a visualization of the data. However, confounding covariates can make patterns related to the scientific question of interest difficult to observe. We provide 'aPCoA' as an easy-to-use tool to improve data visualization in this context, enabling enhanced presentation of the effects of interest. Details are described in Yushu Shi, Liangliang Zhang, Kim-Anh Do, Christine Peterson and Robert Jenq (2020) Bioinformatics, Volume 36, Issue 13, 4099-4101.
License GPL (>= 2)
Depends R (>= 3.5.0)
Imports vegan, randomcoloR, ape, car, cluster
NeedsCompilation no
Repository CRAN
Date/Publication 2021-12-13 08:10:02 UTC

R topics documented:

- aPCoA
- Tasmania

Index
aPCoA

Covariate Adjusted PCoA Plot

Description

Adjusted confounding covariates to show the effect of the primary covariate in a PCoA plot. This method is designed for non-Euclidean distance. This function will plot the original PCoA plot along with the covariate adjusted PCoA plot.

Usage

aPCoA(formula, data, maincov, drawEllipse = TRUE, drawCenter = TRUE, pch = 19, cex = 2, lwd = 3, col = NULL, ...)

Arguments

formula A typical formula such as Y~ A, but here Y is a dissimilarity distance. The formula has the same requirements as in adonis function of the vegan package.
data A dataset with the rownames the same as the rownames in distance. This dataset should include both the confounding covariate and the primary covariate.
maincov the covariate of interest in the dataset, must be a factor
drawEllipse Do you want to draw the 95% confidence ellipse for each cluster?
drawCenter Do you want to show the connection between cluster center (medoid) and cluster members?
pch Point shapes
cex Number indicating the amount by which plotting text and symbols should be scaled relative to the default.
lwd Line width of the ellipses
col Color for plot. If not provided by user, will use default distinct colors
... Arguments passed to ‘dataEllipse’.

Value

Two PCoA plots. One is the original one, while the other is the PCoA plot after adjusting for the confounding covariate.

plotMatrix The matrix for plotting the adjusted PCoA plot.

References

Examples

```r
library(vegan)
library(aPCoA)
data("Tasmania")
data<-data.frame(treatment=Tasmania$treatment,block=Tasmania$block)
bray<-vegdist(Tasmania$abund, method="bray")
rownames(data)<-rownames(as.matrix(bray))
opar<-par(mfrow=c(1,2),
  mar=c(3.1, 3.1, 3.1, 5.1),
  mgp=c(2, 0.5, 0),
  oma=c(0, 0, 0, 4))
result<-aPCoA(bray~block,data,treatment)
par(opar)
```

Tasmania Dataset

Description

A classic dataset used in the R package mvabund. This is a study on the effects of disturbance from a soldier crab on 56 species of meiobenthos, which are small invertebrates. Eight of the sixteen observations in the data set correspond to crab disturbances. Besides the crab disturbance, there are also four different locations in the study design, where observations from each location are comprised of two disturbed and two undisturbed ones.

Usage

```r
data(Tasmania)
```

Format

- **abund**: The counts of 56 Meibenthos species.
- **treatment**: This observation has been disturbed by a crab or not.
- **block**: The location indicator of the observation.

References


Examples

```r
data("Tasmania")
```
Index

aPCoA, 2
Tasmania, 3