graph.rdaTest R Documentation

Drawing RDA triplots

Description

This function produces triplots from the output list of the RDA function 'rdaTest'.

Usage

```
graph.rdaTest(
```

rdaTest.out, xax=1, yax=2, scaling=1, graph.type="notchosen", plot.sites=TRUE, plot.spc=TRUE, plot.env=TRUE, binary=NULL, height=6, width=6, xlim=NULL, ylim=NULL, ell=NULL, alpha=0.95, ell.axis=FALSE, mul.spc=0.8, mul.env=0.90, mul.text=0.10, pos.ell=NULL, pos.bin=NULL, pos.sites=1, label.sites=TRUE, label.spc=TRUE, label.env=TRUE, label.ell=TRUE, col.env="blue", col.spc="red", col.site="black", col.ell="black", lty.env=1, lty.spc=1, lty.ell=1, lty.axis=2, cex=1, cex.lab=1, cex.axis=1, lwd=1, len=0.1, saveplot=FALSE, path=NULL, mar.perc=NULL, select.spc=NULL)

Arguments

rdaTest.out name of the rdaTest output object. The most simple call to this function is to

provide only the 'rdaTest.out' output object name. Ex.: graph.rdaTest(toto).

xax, yax canonical axes that will be used for the triplot. These value can be negative; if so,

the plot axes will be reversed.

scaling allows the user to choose between the two scaling types, 1 or 2, for the triplots:

scaling = 1: distance triplot (default);

scaling = 2: correlation triplot.

graph.type either "F" or "Z" -- "F"= site scores, "Z" = fitted site scores.

plot.sites if TRUE, the sites will be plotted as small circles.

plot.spc if TRUE, the species (or other response variables) will be plotted.
plot.env if TRUE, the explanatory (environmental) variables will be plotted.
a vector specifying which of the environmental variables are binary; the

corresponding data points will be averaged and represented by a symbol instead of

an arrow, as in Canoco.

• if set to FALSE, the graph will only appear in a window (quartz);

• if set to TRUE, no graph will be plotted on the screen; a pdf file will created.

path complete path to the file in which the plot will be saved; example:

"~/Desktop/toto.pdf"

width, height size of the triplot in inches.

xlim, ylim vectors describing the minimum and maximum values of the plotted region.

lwd line width for the axes, arrows, and ellipses.

len length of the arrow heads.

mar.perc allows to increase or decrease the margin size, using a percentage of the total

width and height of the graph; the value must be between 0 and 1.

select.spc vector containing a selection of the species, to be drawn in the biplot. Draw all

species if select.spc=NULL (default value).

Parameters controlling the confidence ellipses around groups of points

ell draw confidence ellipses around groups of points according to a vector defining

the group assignments of the objects; see section **Examples** below.

alpha confidence region of the ellipses (e.g. 0.90, 0.80, etc.) Default value: 0.95. lty.ell drawing type for ellipses; see 'lty' in the help documentation file of 'par'. the ellipses and group identification text color; see 'lty' in the help file of 'par'.

ell.axis if TRUE, draw the major and minor axes of the ellipses.

pos.ell offset the group names relative to the ellipse centers: 1=bottom, 2=left, 3=top,

4=right.

Parameters for arrows and text

label.sites if TRUE, labels are added to the site symbols. label.spc if TRUE, labels are added to the species arrows.

label.env if TRUE, labels are added to the environmental variable arrows.

label.ell if TRUE, labels (numbers) are added to the group ellipses.

col.env color for the environmental variable arrows and text.

col.spc color for the species arrows and text. col.site color for the site points and text.

lty.env environmental variable arrows drawing type; see 'lty' in the help documentation

file of 'par'.

lty.spc species arrows drawing type; see 'lty' in the help documentation file of 'par'.

lty.axis drawing type for axes, full lines, dotted lines, etc. cex.xxx change of the font size, with respect to a standard of 1.

mul.spc multiplier based on the range of the plot axes, which changes to the length of the

species arrows proportionally; 1 = maximum relative length in the set of arrows.

mul.env same, for the explanatory variables

mul.text text position at the end of each arrow plus a percentage; default is 10%. pos.bin offset the names of the binary variables: 1=bottom, 2=left, 3=top, 4=right.

pos.sites offset the site labels: 1=bottom, 2=left, 3=top, 4=right.

Details

A triplot of the results of the canonical redundancy analysis (RDA) is drawn, using either scaling 1 or scaling 2.

Scaling type 1 triplot uses matrices Z or F for objects and U for variables; notation as in Legendre and Legendre (1998, Section 11.1.1). This projection preserves the Euclidean distances among the objects.

Scaling type 2 triplot uses matrices ZSc2 or FSc2 for objects and USc2 for variables. This projection preserves the correlations among the response variables' fitted values.

Value

The function plots a canonical ordination triplot to a graphics window or a PDF file.

Reference

Legendre, P. and Legendre, L. 1998. Numerical Ecology. 2nd English ed. Elsevier, Amsterdam.

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See Also

rdaTest

Examples

```
## Example from Legendre and Legendre (1998), p. 590, Table 11.3
Table11.3 = read.table(file.choose())
                                            # File "Table 11-3.txt"
Y = Table 11.3[,1:6]
X = Table 11.3[,7:10]
result = rdaTest(Y, X, testF=TRUE, nperm=999)
graph.rdaTest(result, graph.type="Z")
## Draw 3 species only in the biplot
vec = c(1,3,6)
graph.rdaTest(result, graph.type="Z", select.spc=vec)
## Reverse the orientation of the abscissa ('xax') in the plot
graph.rdaTest(result, xax=-1, yax=2, mul.spc=0.90, mul.env=0.70, mul.text=0.10, scaling=1,
graph.type="F", mai.perc=0.15, pos.sites=3)
## Draw confidence ellipses around groups of points, spider data
spiders.spe = read.table("Spiders 28x12 spe.txt")
spiders.hel = decostand(spiders.spe, "hellinger")
spiders.env = read.table("Spiders 28x4 env.txt")
result.spiders = rdaTest(spiders.hel, spiders.env)
# 'vec' divides the sites into three a priori group
graph.rdaTest(result.spiders, graph.type="Z", ell=vec, lty.ell=3, pos.sites=4, mar.perc=0.15)
## Plot the sites and environmental variables, but not the species
graph.rdaTest(result.spiders, graph.type="Z", plot.spc=FALSE, ell=vec, lty.ell=3, pos.sites=4,
mar.perc=0.15)
```