Package 'AgroR'

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Type Package

Title Experimental Statistics and Graphics for Agricultural Sciences

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Maintainer Gabriel Danilo Shimizu <gabrield.shimizu@gmail.com>

Description Performs the analysis of completely randomized experimental designs (CRD), randomized blocks (RBD) and Latin square (LSD), experiments in double and triple factorial scheme (in CRD and RBD), experiments in subdivided plot scheme (in CRD and RBD), subdivided and joint analysis of experiments in CRD and RBD, linear regression analysis, test for two samples. The package performs analysis of variance, ANOVA assumptions and multiple comparison test of means or regression, according to Pimentel-Gomes (2009, ISBN: 978-85-7133-055-9), nonparametric test (Conover, 1999, ISBN: 0471160687), test for two samples, joint analysis of experiments according to Ferreira (2018, ISBN: 978-85-7269-566-4) and generalized linear model (glm) for binomial and Poisson family in CRD and RBD (Carvalho, FJ (2019), <doi:10.14393/ufu.te.2019.1244>). It can also be used to obtain descriptive measures and graphics, in addition to correlations and creative graphics used in agricultural sciences (Agronomy, Zootechnics, Food Science and related areas).

NeedsCompilation no

Author Gabriel Danilo Shimizu [aut, cre]
(https://orcid.org/0000-0001-8524-508X>https://orcid.org/0000-0003-2778-8654>https://orcid.org/0000-0003-2778-8654>https://orcid.org/0000-0001-9700-9375>https://orcid.org/

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ааср

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Utils: Area under the curve

Description

Performs the calculation of the area under the progress curve. Initially created for the plant disease area, whose name is "area under the disease progress curve", it can be adapted to various areas of agrarian science.

Usage

aacp(data)

Arguments

data

Data.frame containing evaluations in columns. Column names must be numeric and not dates or characters

Value

Returns a vector with the area values under the curve

Note

Just enter the data. Exclude treatment columns. See example.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

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References

Campbell, C. L., and Madden, L. V. (1990). Introduction to plant disease epidemiology. John Wiley and Sons.

See Also

transf, sketch

Examples

```
# Using the simulate1 dataset
data("simulate1")
# Converting to readable format for function
dados=cbind(simulate1[simulate1$tempo==1,3],
           simulate1[simulate1$tempo==2,3],
           simulate1[simulate1$tempo==3,3],
           simulate1[simulate1$tempo==4,3],
           simulate1[simulate1$tempo==5,3],
           simulate1[simulate1$tempo==6,3])
colnames(dados)=c(1,2,3,4,5,6)
dados
# Creating the treatment vector
resp=aacp(dados)
trat=simulate1$trat[simulate1$tempo==1]
# Analyzing by DIC function
DIC(trat,resp)
```

aristolochia

Dataset: Germination of seeds of Aristolochia sp. as a function of temperature.

Description

The data come from an experiment conducted at the Seed Analysis Laboratory of the Agricultural Sciences Center of the State University of Londrina, in which five temperatures (15, 20, 25, 30 and 35C) were evaluated in the germination of *Aristolochia elegans*. The experiment was conducted in a completely randomized design with four replications of 25 seeds each.

Usage

```
data("aristolochia")
```

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Format

```
data.frame containing data set
trat numeric vector with factor 1
resp Numeric vector with response
```

See Also

cloro, laranja, enxofre, laranja, mirtilo, passiflora, phao, porco, pomegranate, simulate1, simulate2, simulate3, tomate, weather

Examples

```
data(aristolochia)
```

barfacet

Graph: Bar graph for one factor with facets

Description

This is a function of the bar graph for one factor with facets

Usage

```
barfacet(
  model,
  facet = NULL,
  theme = theme_bw(),
  horiz = FALSE,
  geom = "bar",
  fill = "lightblue",
  pointsize = 4.5,
  width.bar = 0.15,
  facet.background = "gray80"
)
```

Arguments

```
DIC, DBC or DQL object
model
facet
                  vector with facets
                  ggplot2 theme
theme
horiz
                  horizontal bar or point (default is FALSE)
                  graph type (columns or segments)
geom
fill
                  fill bars
                  Point size
pointsize
width.bar
                  width of the error bars of a regression graph.
facet.background
```

Color background in facet

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Value

Returns a bar chart for one factor

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

Examples

bargraph_onefactor

Graph: Group DIC, DBC and DQL functions column charts

Description

Groups two or more column charts exported from DIC, DBC or DQL function

Usage

```
bargraph_onefactor(
   analysis,
   labels = NULL,
   ocult.facet = FALSE,
   ocult.box = FALSE,
   facet.size = 14,
   ylab = NULL,
   width.bar = 0.3,
   width.col = 0.9,
   sup = NULL
)
```

```
analysis List with DIC, DBC or DQL object labels Vector with the name of the facets ocult.facet Hide facets
ocult.box Hide box
```

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facet.size	Font size facets
ylab	Y-axis name
width.bar	Width error bar
width.col	Width Column

sup Number of units above the standard deviation or average bar on the graph

Value

Returns a column chart grouped by facets

Examples

```
library(AgroR)
data("laranja")
a=with(laranja, DBC(trat, bloco, resp, ylab = "Number of fruits/plants"))
b=with(laranja, DBC(trat, bloco, resp, ylab = "Number of fruits/plants"))
c=with(laranja, DBC(trat, bloco, resp, ylab = "Number of fruits/plants"))
bargraph_onefactor(analysis = list(a,b,c), labels = c("One", "Two", "Three"),ocult.box = TRUE)
```

bargraph_twofactor

Graph: Group FAT2DIC, FAT2DBC, PSUBDIC or PSUBDBC functions column charts

Description

Groups two or more column charts exported from FAT2DIC, FAT2DBC, PSUBDIC or PSUBDBC function

Usage

```
bargraph_twofactor(
  analysis,
  labels = NULL,
  ocult.facet = FALSE,
  ocult.box = FALSE,
  facet.size = 14,
  ylab = NULL,
  width.bar = 0.3,
  sup = NULL
)
```

Arguments

analysis List with DIC, DBC or DQL object labels Vector with the name of the facets

ocult.facet Hide facets

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ocult.box	Hide box
facet.size	Font size facets
ylab	Y-axis name
width.bar	Width bar

sup Number of units above the standard deviation or average bar on the graph

Value

Returns a column chart grouped by facets

Examples

```
library(AgroR)
data(corn)
a=with(corn, FAT2DIC(A, B, Resp, quali=c(TRUE, TRUE),ylab="Heigth (cm)"))
b=with(corn, FAT2DIC(A, B, Resp, mcomp="sk", quali=c(TRUE, TRUE),ylab="Heigth (cm)"))
bargraph_twofactor(analysis = list(a,b), labels = c("One","Two"),ocult.box = TRUE)
```

barplot_positive

Graph: Positive barplot

Description

Column chart with two variables that assume a positive response and represented by opposite sides, such as dry mass of the area and dry mass of the root

Usage

```
barplot_positive(
   a,
   b,
   ylab = "Response",
   var_name = c("Var1", "Var2"),
   legend.title = "Variable",
   fill_color = c("darkgreen", "brown"),
   width.col = 0.9,
   width.bar = 0.2
)
```

```
a Object of DIC, DBC or DQL functions
b Object of DIC, DBC or DQL functions
ylab Y axis names
var_name Name of the variable
legend.title Legend title
```

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```
fill_color Bar fill color
width.col Width Column
width.bar Width error bar
```

Value

The function returns a column chart with two positive sides

Note

When there is only an effect of the isolated factor in the case of factorial or subdivided plots, it is possible to use the barplot_positive function.

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
```

See Also

```
radargraph, sk_graph, plot_TH, corgraph, spider_graph, line_plot
```

Examples

```
data("passiflora")
attach(passiflora)
a=with(passiflora, DBC(trat, bloco, MSPA))
b=with(passiflora, DBC(trat, bloco, MSR))
barplot_positive(a, b, var_name = c("DMAP","DRM"), ylab = "Dry root (g)")
a=with(passiflora, DIC(trat, MSPA,test = "noparametric"))
b=with(passiflora, DIC(trat, MSR))
barplot_positive(a, b, var_name = c("DMAP","DRM"), ylab = "Dry root (g)")
```

bar_dunnett

Graph: Barplot for Dunnett test

Description

The function performs the construction of a column chart of Dunnett's test.

Usage

```
bar_dunnett(
  output.dunnett,
  ylab = "Response",
  xlab = "",
  fill = c("#F8766D", "#00BFC4"),
  sup = NA,
```

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```
add.mean = TRUE,
round = 2
)
```

Arguments

$\verb"output.dunnett"$	Numerical or complex vector with treatments
ylab	Variable response name (Accepts the expression() function)
xlab	Treatments name (Accepts the expression() function)
fill	$Fill\ column.\ Use\ vector\ with\ two\ elements\ c(control,\ different\ treatment)$
sup	Number of units above the standard deviation or average bar on the graph
add.mean	Plot the average value on the graph (default is TRUE)
round	Number of cells

Value

Returns a column chart of Dunnett's test. The colors indicate difference from the control.

Examples

```
# randomized block design in factorial double
library(AgroR)
data(cloro)
attach(cloro)
respAd=c(268, 322, 275, 350, 320)
a=FAT2DBC.ad(f1, f2, bloco, resp, respAd,
          ylab="Number of nodules",
          legend = "Stages",mcomp="sk")
data=rbind(data.frame(trat=paste(f1,f2,sep = ""),bloco=bloco,resp=resp),
         data.frame(trat=c("Test","Test","Test","Test","Test"),
                  bloco=unique(bloco),resp=respAd))
a= with(data,dunnett(trat = trat,
              resp = resp,
              control = "Test",
              block=bloco,model = "DBC"))
bar_dunnett(a)
```

bar_graph

Graph: Bar graph for one factor

Description

This is a function of the bar graph for one factor

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Usage

```
bar_graph(
  model,
  fill = "lightblue",
  horiz = TRUE,
  width.col = 0.9,
  axis.0 = FALSE
)
```

Arguments

model	DIC, DBC or DQL object
fill	fill bars
horiz	Horizontal Column (default is TRUE)
width.col	Width Column
axis.0	If TRUE causes the columns or bars to start just above the axis line.

Value

Returns a bar chart for one factor

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

See Also

```
radargraph, barplot_positive, plot_TH, plot_TH1, corgraph, spider_graph, line_plot, plot_cor, plot_interaction, plot_jitter, seg_graph, TBARPLOT.reverse
```

```
data("laranja")
a=with(laranja, DBC(trat, bloco, resp,
    mcomp = "sk",angle=45,
    ylab = "Number of fruits/plants"))
bar_graph(a,horiz = FALSE)
```

bar_graph2

bar_graph2

Graph: Bar graph for one factor model 2

Description

This is a function of the bar graph for one factor

Usage

```
bar_graph2(
 model,
 point.color = "black",
 point.size = 2,
 point.shape = 16,
  text.color = "black",
  label.color = "black",
 bar.color = "black",
  title.size = 14,
 y.text = 0,
  add.info = NA,
 y.info = 0,
 width.col = 0.9,
 width.bar = 0,
  color.info = "black",
  fill = "lightblue"
)
```

model	DIC, DBC or DQL object
point.color	Point color
point.size	Point size
point.shape	Format point
text.color	Text color
label.color	Label color
bar.color	Errorbar color
title.size	Title size
y.text	Y-axis height for x-axis legend
add.info	Add other information
y.info	Y-axis height for other information
width.col	Width Column
width.bar	Width error bar
color.info	Color text information
fill	Fill bars

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Value

Returns a bar chart for one factor

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

See Also

radargraph, barplot_positive, plot_TH, plot_TH1, corgraph, spider_graph, line_plot, plot_cor, plot_interaction, plot_jitter, seg_graph, TBARPLOT.reverse

Examples

bean

Dataset: Bean

Description

An experiment to evaluate the effect of different strains of Azospirillum on common bean cultivar IPR Sabia was carried out in a greenhouse. A completely randomized design with five strains was used. of Azospirillum (treatments) and five repetitions. The response variable analyzed was grain production per plant (g plant-1).

Usage

```
data("bean")
```

Format

```
data.frame containing data set

trat numeric vector with treatment

prod Numeric vector with grain production per plant
```

See Also

aristolochia, cloro, laranja, enxofre, laranja, mirtilo, passiflora, phao, porco, pomegranate, simulate1, simulate2, simulate3, tomate, weather

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Examples

data(bean)

cloro

Dataset: Sodium dichloroisocyanurate in soybean

Description

An experiment was conducted in a greenhouse in pots at the State University of Londrina. The work has the objective of evaluating the application of sodium dichloroisocyanurate (DUP) in soybean in 4 periods of application in soybean inoculated or not with Rhizobium and its influence on the number of nodules. The experiment was conducted in a completely randomized design with five replications.

Usage

data(cloro)

Format

data.frame containing data set

f1 Categorical vector with factor 1

f2 Categorical vector with factor 2

bloco Categorical vector with block

resp Numeric vector with number nodules

References

Rony Kauling Tonelli. Efeito do uso de dicloroisocianurato de sodio sobre a nodulacao em raizes de soja. 2016. Trabalho de Conclusao de Curso. (Graduacao em Agronomia) - Universidade Estadual de Londrina.

See Also

enxofre, laranja, mirtilo, pomegranate, porco, sensorial, simulate1, simulate2, simulate3, tomate, weather, phao, passiflora, aristolochia

Examples

data(cloro)

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Utils: Interval of confidence for groups

Description

Calculates confidence interval for groups

Usage

```
confinterval(resp, group, alpha = 0.95, type = "upper")
```

Arguments

resp	numeric vector with	responses
1 636	mumeric vector with	responses

group vector with groups or list with two factors

alpha confidence level of the interval

type lower or upper range

Value

returns a numeric vector with confidence interval grouped by treatment.

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conjdbc

Analysis: Joint analysis of experiments in randomized block design

Description

Function of the AgroR package for joint analysis of experiments conducted in a randomized qualitative or quantitative single-block design with balanced data.

Usage

```
conjdbc(
  trat,
  block,
  local,
  response,
  transf = 1,
  constant = 0,
  norm = "sw",
  homog = "bt",
  homog.value = 7,
  theme = theme_classic(),
 mcomp = "tukey",
  quali = TRUE,
  alpha.f = 0.05,
  alpha.t = 0.05,
  grau = NA,
 ylab = "response",
 title = "",
xlab = "",
  fill = "lightblue",
  angulo = 0,
  textsize = 12,
  dec = 3,
  family = "sans",
  errorbar = TRUE
)
```

trat	Numerical or complex vector with treatments
block	Numerical or complex vector with blocks
local	Numeric or complex vector with locations or times
response	Numerical vector containing the response of the experiment.
transf	Applies data transformation (default is 1; for log consider 0)
constant	Add a constant for transformation (enter value)

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norm Error normality test (*default* is Shapiro-Wilk) homog Homogeneity test of variances (*default* is Bartlett)

homog.value Reference value for homogeneity of experiments. By default, this ratio should

not be greater than 7

theme ggplot2 theme (default is theme_classic())

mcomp Multiple comparison test (Tukey (default), LSD, Scott-Knott and Duncan)
quali Defines whether the factor is quantitative or qualitative (default is qualitative)

alpha.f Level of significance of the F test (*default* is 0.05)

alpha.t Significance level of the multiple comparison test (default is 0.05) grau Degree of polynomial in case of quantitative factor (default is 1) ylab Variable response name (Accepts the expression() function)

title Graph title

xlab Treatments name (Accepts the *expression*() function)

fill Defines chart color (to generate different colors for different treatments, define

fill = "trat"

angulo x-axis scale text rotation

textsize Font size

dec Number of cells family Font family

errorbar Plot the standard deviation bar on the graph (In the case of a segment and column

graph) - default is TRUE

Value

Returns the assumptions of the analysis of variance, the assumption of the joint analysis by means of a QMres ratio matrix, the analysis of variance, the multiple comparison test or regression.

Note

In this function there are three possible outcomes. When the ratio between the experiments is greater than 7, the separate analyzes are returned, without however using the square of the joint residue. When the ratio is less than 7, but with significant interaction, the effects are tested using the square of the joint residual. When there is no significant interaction and the ratio is less than 7, the joint analysis between the experiments is returned.

The ordering of the graph is according to the sequence in which the factor levels are arranged in the data sheet. The bars of the column and segment graphs are standard deviation.

In the final output when transformation (transf argument) is different from 1, the columns resp and respo in the mean test are returned, indicating transformed and non-transformed mean, respectively.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi

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References

Ferreira, P. V. Estatistica experimental aplicada a agronomia. Edufal, 2018.

Principles and procedures of statistics a biometrical approach Steel, Torry and Dickey. Third Edition 1997

Multiple comparisons theory and methods. Departament of statistics the Ohio State University. USA, 1996. Jason C. Hsu. Chapman Hall/CRC.

Practical Nonparametrics Statistics. W.J. Conover, 1999

Ramalho M.A.P., Ferreira D.F., Oliveira A.C. 2000. Experimentacao em Genetica e Melhoramento de Plantas. Editora UFLA.

Scott R.J., Knott M. 1974. A cluster analysis method for grouping mans in the analysis of variance. Biometrics, 30, 507-512.

Examples

conjdic

Analysis: Joint analysis of experiments in completely randomized design

Description

Function of the AgroR package for joint analysis of experiments conducted in a completely randomized design with a qualitative or quantitative factor with balanced data.

Usage

```
conjdic(
  trat,
  repet,
  local,
  response,
  transf = 1,
  constant = 0,
```

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```
norm = "sw",
homog = "bt",
mcomp = "tukey",
homog.value = 7,
quali = TRUE,
alpha.f = 0.05,
alpha.t = 0.05,
grau = NA,
theme = theme_classic(),
ylab = "response",
title = "",
xlab = "",
color = "rainbow",
fill = "lightblue",
angulo = 0,
textsize = 12,
dec = 3,
family = "sans",
errorbar = TRUE
```

trat	Numerical or complex vector with treatments
repet	Numerical or complex vector with repetitions
local	Numeric or complex vector with locations or times
response	Numerical vector containing the response of the experiment.
transf	Applies data transformation (default is 1; for log consider 0)
constant	Add a constant for transformation (enter value)
norm	Error normality test (default is Shapiro-Wilk)
homog	Homogeneity test of variances (default is Bartlett)
mcomp	Multiple comparison test (Tukey (default), LSD, Scott-Knott and Duncan)
homog.value	Reference value for homogeneity of experiments. By default, this ratio should not be greater than 7
quali	Defines whether the factor is quantitative or qualitative (<i>default</i> is qualitative)
alpha.f	Level of significance of the F test (default is 0.05)
alpha.t	Significance level of the multiple comparison test (default is 0.05)
grau	Degree of polynomial in case of quantitative factor (default is 1)
theme	ggplot2 theme (default is theme_classic())
ylab	Variable response name (Accepts the expression() function)
title	Graph title
xlab	Treatments name (Accepts the expression() function)
color	When the columns are different colors (Set fill-in argument as "trat")

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fill Defines chart color (to generate different colors for different treatments, define

fill = "trat")

angulo x-axis scale text rotation

textsize Font size

dec Number of cells family Font family

errorbar Plot the standard deviation bar on the graph (In the case of a segment and column

graph) - default is TRUE

Value

Returns the assumptions of the analysis of variance, the assumption of the joint analysis by means of a QMres ratio matrix, the analysis of variance, the multiple comparison test or regression.

Note

In this function there are three possible outcomes. When the ratio between the experiments is greater than 7, the separate analyzes are returned, without however using the square of the joint residue. When the ratio is less than 7, but with significant interaction, the effects are tested using the square of the joint residual. When there is no significant interaction and the ratio is less than 7, the joint analysis between the experiments is returned.

The ordering of the graph is according to the sequence in which the factor levels are arranged in the data sheet. The bars of the column and segment graphs are standard deviation.

In the final output when transformation (transf argument) is different from 1, the columns resp and respo in the mean test are returned, indicating transformed and non-transformed mean, respectively.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Leandro Simoes Azeredo Goncalves

Rodrigo Yudi Palhaci Marubayashi

References

Ferreira, P. V. Estatistica experimental aplicada a agronomia. Edufal, 2018.

Principles and procedures of statistics a biometrical approach Steel, Torry and Dickey. Third Edition 1997

Multiple comparisons theory and methods. Departament of statistics the Ohio State University. USA, 1996. Jason C. Hsu. Chapman Hall/CRC.

Practical Nonparametrics Statistics. W.J. Conover, 1999

Ramalho M.A.P., Ferreira D.F., Oliveira A.C. 2000. Experimentacao em Genetica e Melhoramento de Plantas. Editora UFLA.

Scott R.J., Knott M. 1974. A cluster analysis method for grouping mans in the analysis of variance. Biometrics, 30, 507-512.

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Examples

```
library(AgroR)
data(mirtilo)
with(mirtilo, conjdic(trat, bloco, exp, resp))
```

conjfat2dbc

Analysis: Joint analysis of experiments in randomized block design in scheme factorial double

Description

Function of the AgroR package for joint analysis of experiments conducted in a randomized factorial double in block design with balanced data. The function generates the joint analysis through two models. Model 1: F-test of the effects of Factor 1, Factor 2 and F1 x F2 interaction are used in reference to the mean square of the interaction with the year. Model 2: F-test of the Factor 1, Factor 2 and F1 x F2 interaction effects are used in reference to the mean square of the residual.

Usage

```
conjfat2dbc(
  f1,
  f2,
  block,
  experiment,
  response,
  transf = 1,
  constant = 0,
  model = 1,
  norm = "sw",
  homog = "bt",
  homog.value = 7,
  alpha.f = 0.05,
  alpha.t = 0.05
```

Arguments

f1	Numeric or complex vector with factor 1 levels
f2	Numeric or complex vector with factor 2 levels
block	Numerical or complex vector with blocks
experiment	Numeric or complex vector with locations or times

response Numerical vector containing the response of the experiment. transf Applies data transformation (default is 1; for log consider 0)

constant Add a constant for transformation (enter value)
model Define model of the analysis of variance

conjfat2dbc 23

norm	Error normality test (<i>default</i> is Shapiro-Wilk)
homog	Homogeneity test of variances (default is Bartlett)
homog.value	Reference value for homogeneity of experiments. By default, this ratio should not be greater than 7
alpha.f	Level of significance of the F test (default is 0.05)
alpha.t	Significance level of the multiple comparison test (default is 0.05)

Value

Returns the assumptions of the analysis of variance, the assumption of the joint analysis by means of a QMres ratio matrix and analysis of variance

Note

The function is still limited to analysis of variance and assumptions only.

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

References

Ferreira, P. V. Estatistica experimental aplicada a agronomia. Edufal, 2018.

Principles and procedures of statistics a biometrical approach Steel, Torry and Dickey. Third Edition 1997

Multiple comparisons theory and methods. Departament of statistics the Ohio State University. USA, 1996. Jason C. Hsu. Chapman Hall/CRC.

Practical Nonparametrics Statistics. W.J. Conover, 1999

Ramalho M.A.P., Ferreira D.F., Oliveira A.C. 2000. Experimentacao em Genetica e Melhoramento de Plantas. Editora UFLA.

```
library(AgroR)
ano=factor(rep(c(2018,2019,2020),e=48))
f1=rep(rep(c("A","B","C"),e=16),3)
f2=rep(rep(rep(c("a1","a2","a3","a4"),e=4),3),3)
resp=rnorm(48*3,10,1)
bloco=rep(c("b1","b2","b3","b4"),36)
dados=data.frame(ano,f1,f2,resp,bloco)
with(dados,conjfat2dbc(f1,f2,bloco,ano,resp, model=1))
```

24 corgraph

corgraph

Graph: Correlogram

Description

Correlation analysis function (Pearson or Spearman)

Usage

```
corgraph(
  data,
  axissize = 12,
  legendsize = 12,
  legendposition = c(0.9, 0.2),
  legendtitle = "Correlation",
  method = "pearson",
  pallete = "RdBu",
  color.marginal = "gray50",
  size.tile.lty = 1,
  size.label.cor = 1,
  fill.label.cor = "lightyellow",
  font.family = "sans"
)
```

Arguments

```
data
                  data.frame with responses
axissize
                  Axes font size (default is 12)
                  Legend font size (default is 12)
legendsize
legendposition Legend position (default is c(0.9,0.2))
legendtitle
                  Legend title (default is "Correlation")
method
                  Method correlation (default is Pearson)
pallete
                  If a string, will use that named palette. See scale_fill_distiller in the ggplot2.
color.marginal Box border color
                  Box margin line thickness
size.tile.lty
size.label.cor Label font size
fill.label.cor Label fill color
font.family
                  Font family (default is sans)
```

Value

The function returns a correlation matrix

corn 25

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br> Leandro Simoes Azeredo Goncalves Rodrigo Yudi Palhaci Marubayashi

Examples

```
data("pomegranate")
corgraph(pomegranate[,-1])
```

corn

Dataset: Corn

Description

A 3 x 2 factorial experiment was carried out to compare three new corn hybrids considering the change in sowing density, being 55 thousand or 65 thousand seeds per hectare. For this case, the researcher is not interested in estimating values for other densities, but only in verifying if one density differs from the other. The experiment was carried out according to a completely randomized design with 4 repetitions of each treatment.

Usage

data(corn)

Format

data.frame containing data set

- A Categorical vector with hybrids
- B Categorical vector with density

resp Numeric vector with response

See Also

enxofre, laranja, mirtilo, pomegranate, porco, sensorial, simulate1, simulate2, simulate3, tomate, weather, phao, passiflora, aristolochia

Examples

data(corn)

26 cor_ic

cor_ic

Graph: Plot Pearson correlation with interval of confidence

Description

Plot Pearson correlation with interval of confidence

Usage

```
cor_ic(
  data,
  background = TRUE,
  axis.size = 12,
  ylab = "",
  xlab = "Correlation (r)",
  theme = theme_classic()
)
```

Arguments

data	data.frame with responses
background	background fill (default is TRUE)
axis.size	Axes font size (default is 12)
ylab	Variable response name (Accepts the <i>expression</i> () function)
xlab	Treatments name (Accepts the expression() function)
theme	ggplot theme (default is theme_classic())

Value

The function returns a new graphical approach to correlation.

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

```
data("pomegranate")
cor_ic(pomegranate[,-1])
```

covercrops 27

covercrops

Dataset: Covercrops

Description

Consider a 3 ×3 factorial experiment in randomized blocks, with 4 replications, on the influence of three new soybean cultivars (A1, A2 and A3) and the use of three types of green manure (B1, B2 and B3) on yield in 100 m2 plots.

Usage

data(covercrops)

Format

data.frame containing data set

- A Categorical vector with cultivars
- B Categorical vector with green manure

Bloco Categorical vector with block

Resp Numeric vector with yield

See Also

enxofre, laranja, mirtilo, pomegranate, porco, sensorial, simulate1, simulate2, simulate3, tomate, weather, phao, passiflora, aristolochia

Examples

data(covercrops)

DBC

Analysis: Randomized block design

Description

This is a function of the AgroR package for statistical analysis of experiments conducted in a randomized block and balanced design with a factor considering the fixed model. The function presents the option to use non-parametric method or transform the dataset.

28 DBC

Usage

```
DBC(
  trat,
  block,
  response,
  norm = "sw",
  homog = "bt",
  alpha.f = 0.05,
  alpha.t = 0.05,
  quali = TRUE,
 mcomp = "tukey",
  grau = 1,
  transf = 1,
  constant = 0,
  test = "parametric",
  geom = "bar",
  theme = theme_classic(),
  sup = NA,
  CV = TRUE,
  ylab = "response",
  xlab = "",
  textsize = 12,
  labelsize = 4,
  fill = "lightblue",
  angle = 0,
  family = "sans",
  dec = 3,
 width.column = NULL,
 width.bar = 0.3,
  addmean = TRUE,
  errorbar = TRUE,
  posi = "top",
  point = "mean_sd",
  pointsize = 5,
  angle.label = 0,
 ylim = NA
)
```

trat	Numerical or complex vector with treatments
block	Numerical or complex vector with blocks
response	Numerical vector containing the response of the experiment.
norm	Error normality test (default is Shapiro-Wilk)
homog	Homogeneity test of variances (default is Bartlett)
alpha.f	Level of significance of the F test (<i>default</i> is 0.05)
alpha.t	Significance level of the multiple comparison test (<i>default</i> is 0.05)

DBC 29

quali Defines whether the factor is quantitative or qualitative (*default* is qualitative) mcomp Multiple comparison test (Tukey (*default*), LSD, Scott-Knott and Duncan)

grau Degree of polynomial in case of quantitative factor (*default* is 1)

transf Applies data transformation (default is 1; for log consider 0; 'angular' for angu-

lar transformation)

constant Add a constant for transformation (enter value)

test "parametric" - Parametric test or "noparametric" - non-parametric test

geom graph type (columns, boxes or segments)
theme ggplot2 theme (default is theme_classic())

Sup Number of units above the standard deviation or average bar on the graph CV Plotting the coefficient of variation and p-value of Anova (*default* is TRUE)

ylab Variable response name (Accepts the *expression*() function) xlab Treatments name (Accepts the *expression*() function)

textsize Font size labelsize Label size

fill Defines chart color (to generate different colors for different treatments, define

fill = "trat")

angle x-axis scale text rotation

family Font family dec Number of cells

width.column Width column if geom="bar"

width.bar Width errorbar

addmean Plot the average value on the graph (*default* is TRUE)

errorbar Plot the standard deviation bar on the graph (In the case of a segment and column

graph) - default is TRUE

posi Legend position

point Defines whether to plot mean ("mean"), mean with standard deviation ("mean_sd"

- default) or mean with standard error ("mean_se"). For parametric test it is pos-

sible to plot the square root of QMres (mean_qmres).

pointsize Point size angle.label label angle

ylim Define a numerical sequence referring to the y scale. You can use a vector or the

'seq' command.

Value

The table of analysis of variance, the test of normality of errors (Shapiro-Wilk ("sw"), Lilliefors ("li"), Anderson-Darling ("ad"), Cramer-von Mises ("cvm"), Pearson ("pearson") and Shapiro-Francia ("sf")), the test of homogeneity of variances (Bartlett ("bt") or Levene ("levene")), the test of independence of Durbin-Watson errors, the test of multiple comparisons (Tukey ("tukey"), LSD ("lsd"), Scott-Knott ("sk") or Duncan ("duncan")) or adjustment of regression models up to grade 3 polynomial, in the case of quantitative treatments. Non-parametric analysis can be used by the Friedman test. The column, segment or box chart for qualitative treatments is also returned. The function also returns a standardized residual plot.

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Note

Enable ggplot2 package to change theme argument.

The ordering of the graph is according to the sequence in which the factor levels are arranged in the data sheet. The bars of the column and segment graphs are standard deviation.

CV and p-value of the graph indicate coefficient of variation and p-value of the F test of the analysis of variance.

In the final output when transformation (transf argument) is different from 1, the columns resp and respo in the mean test are returned, indicating transformed and non-transformed mean, respectively.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi

References

Principles and procedures of statistics a biometrical approach Steel, Torry and Dickey. Third Edition 1997

Multiple comparisons theory and methods. Departament of statistics the Ohio State University. USA, 1996. Jason C. Hsu. Chapman Hall/CRC.

Practical Nonparametrics Statistics. W.J. Conover, 1999

Ramalho M.A.P., Ferreira D.F., Oliveira A.C. 2000. Experimentacao em Genetica e Melhoramento de Plantas. Editora UFLA.

Scott R.J., Knott M. 1974. A cluster analysis method for grouping mans in the analysis of variance. Biometrics, 30, 507-512.

Mendiburu, F., and de Mendiburu, M. F. (2019). Package 'agricolae'. R Package, Version, 1-2.

See Also

DIC, DQL

dbc.ad 31

dbc.ad

Analysis: Randomized block design with an additional treatment for quantitative factor

Description

Statistical analysis of experiments conducted in a randomized block design with an additional treatment and balanced design with a factor considering the fixed model.

Usage

```
dbc.ad(
  trat,
 block,
  response,
  responsead,
  grau = 1,
 norm = "sw"
 homog = "bt",
 alpha.f = 0.05,
  theme = theme_classic(),
 ylab = "response",
  xlab = "independent",
  family = "sans",
 posi = "top",
 pointsize = 4.5,
 linesize = 0.8,
 width.bar = NA,
  point = "mean_sd"
)
```

trat	Numerical or complex vector with treatments
block	Numerical or complex vector with blocks
response	Numerical vector containing the response of the experiment.
responsead	Numerical vector with additional treatment responses
grau	Degree of polynomial in case of quantitative factor (<i>default</i> is 1)

32 dbc.ad

norm Error normality test (default is Shapiro-Wilk)
homog Homogeneity test of variances (default is Bartlett)
alpha.f Level of significance of the F test (default is 0.05)

theme ggplot2 theme (default is theme_classic())

ylab Variable response name (Accepts the *expression*() function)

xlab Treatments name (Accepts the *expression*() function)

family Font family
posi Legend position
pointsize Point size

linesize line size (Trendline and Error Bar)

width.bar width of the error bars of a regression graph.

point Defines whether to plot mean ("mean"), mean with standard deviation ("mean sd"

- default) or mean with standard error (default - "mean_se"). For quali=FALSE

or quali=TRUE.

Value

The table of analysis of variance, the test of normality of errors (Shapiro-Wilk ("sw"), Lilliefors ("li"), Anderson-Darling ("ad"), Cramer-von Mises ("cvm"), Pearson ("pearson") and Shapiro-Francia ("sf")), the test of homogeneity of variances (Bartlett ("bt") or Levene ("levene")), the test of independence of Durbin-Watson errors, adjustment of regression models up to grade 3 polynomial. The function also returns a standardized residual plot.

Note

In some experiments, the researcher may study a quantitative factor, such as fertilizer doses, and present a control, such as a reference fertilizer, treated as a qualitative control. In these cases, there is a difference between considering only the residue in the unfolding of the polynomial, removing or not the qualitative treatment, or since a treatment is excluded from the analysis. In this approach, the residue used is also considering the qualitative treatment, a method similar to the factorial scheme with additional control.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br> Leandro Simoes Azeredo Goncalves Rodrigo Yudi Palhaci Marubayashi

```
doses = c(rep(c(1:5),e=3))
resp = c(3, 4, 3, 5, 5, 6, 7, 7, 8, 4, 4, 5, 2, 2, 3)
bloco = rep(c("B1","B2","B3","B4","B5"),3)
dbc.ad(doses, bloco, resp, responsead=rnorm(3,6,0.1),grau=2)
```

DBC.glm

DBC.glm

Analysis: Randomized block design by glm

Description

Statistical analysis of experiments conducted in a randomized block design using a generalized linear model. It performs the deviance analysis and the effect is tested by a chi-square test. Multiple comparisons are adjusted by Tukey.

Usage

```
DBC.glm(
  trat,
 block,
  response,
  glm.family = "binomial",
  quali = TRUE,
  alpha.f = 0.05,
  alpha.t = 0.05,
  geom = "bar",
  theme = theme_classic(),
  sup = NA,
  ylab = "Response",
  xlab = "",
  fill = "lightblue",
  angle = 0,
  family = "sans",
  textsize = 12,
  labelsize = 5,
  dec = 3,
  addmean = TRUE,
  errorbar = TRUE,
  posi = "top",
  point = "mean_sd",
  angle.label = 0
)
```

trat	Numerical or complex vector with treatments
block	Numerical or complex vector with blocks
response	Numerical vector containing the response of the experiment. Use cbind(resp, n-resp) for binomial or quasibinomial family.
glm.family	distribution family considered (default is binomial)
quali	Defines whether the factor is quantitative or qualitative (<i>default</i> is qualitative)

34 DBC.glm

Level of significance of the F test (*default* is 0.05)

alpha.t Significance level of the multiple comparison test (default is 0.05)

geom Graph type (columns, boxes or segments)

theme ggplot2 theme (default is theme_classic())

sup Number of units above the standard deviation or average bar on the graph

ylab Variable response name (Accepts the expression() function)

xlab Treatments name (Accepts the expression() function)

fill Defines chart color (to generate different colors for different treatments, define

fill = "trat")

angle x-axis scale text rotation

family Font family textsize Font size labelsize Label size

alpha.f

dec Number of cells

addmean Plot the average value on the graph (*default* is TRUE)

errorbar Plot the standard deviation bar on the graph (In the case of a segment and column

graph) - default is TRUE

posi Legend position

point Defines whether to plot mean ("mean"), mean with standard deviation ("mean_sd"

- default) or mean with standard error (default - "mean_se").

angle.label label angle

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Leandro Simoes Azeredo Goncalves

Rodrigo Yudi Palhaci Marubayashi

```
data("aristolochia")
attach(aristolochia)
# Assuming the same aristolochia data set, but considering randomized blocks
bloco=rep(paste("B",1:16),5)
resp=resp/2
DBC.glm(trat,bloco, cbind(resp,50-resp), glm.family="binomial")
```

DBCT 35

DBCT

Analysis: Randomized block design evaluated over time

Description

Function of the AgroR package for analysis of experiments conducted in a balanced qualitative, single-factorial randomized block design with multiple assessments over time, however without considering time as a factor.

Usage

```
DBCT(
  trat,
  block,
  time,
  response,
  alpha.f = 0.05,
  alpha.t = 0.05,
 mcomp = "tukey",
  geom = "bar",
  theme = theme_classic(),
  fill = "gray",
  ylab = "Response",
  xlab = "Independent",
  textsize = 12,
  labelsize = 5,
  pointsize = 4.5,
  error = TRUE,
  family = "sans",
  sup = 0,
  addmean = FALSE,
  posi = c(0.1, 0.8),
  legend = "Legend",
  ylim = NA,
  width.bar = 0.2,
  size.bar = 0.8,
  dec = 3,
  xnumeric = FALSE,
  all.letters = FALSE
)
```

Arguments

trat Numerical or complex vector with treatments
block Numerical or complex vector with blocks
time Numerical or complex vector with times

36 DBCT

response Numerical vector containing the response of the experiment.

alpha.f Level of significance of the F test (*default* is 0.05)

alpha.t Significance level of the multiple comparison test (*default* is 0.05)

mcomp Multiple comparison test (Tukey (default), LSD ("lsd"), Scott-Knott ("sk"), Dun-

can ("duncan") and Friedman ("fd"))

geom Graph type (columns - "bar" or segments "point")

theme ggplot2 theme (*default* is theme_classic())

fill Defines chart color (to generate different colors for different treatments, define

fill = "trat"

ylab Variable response name (Accepts the *expression*() function)

xlab Treatments name (Accepts the *expression*() function)

textsize Font size of the texts and titles of the axes

labelsize Font size of the labels

pointsize Point size

error Add error bar (SD)

family Font family

sup Number of units above the standard deviation or average bar on the graph

addmean Plot the average value on the graph (*default* is TRUE)

posi Legend position legend Legend title

ylim Define a numerical sequence referring to the y scale. You can use a vector or the

'seq' command.

width.bar width error bar size.bar size error bar dec Number of cells

xnumeric Declare x as numeric (*default* is FALSE)

all.letters Adds all label letters regardless of whether it is significant or not.

Details

The p-value of the analysis of variance, the normality test for Shapiro-Wilk errors, the Bartlett homogeneity test of variances, the independence of Durbin-Watson errors and the multiple comparison test (Tukey, Scott-Knott, LSD or Duncan).

Value

The function returns the p-value of Anova, the assumptions of normality of errors, homogeneity of variances and independence of errors, multiple comparison test, as well as a line graph

Note

The ordering of the graph is according to the sequence in which the factor levels are arranged in the data sheet. The bars of the column and segment graphs are standard deviation.

desc 37

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Leandro Simoes Azeredo Gonçalves

Rodrigo Yudi Palhaci Marubayashi

References

Principles and procedures of statistics a biometrical approach Steel & Torry & Dickey. Third Edition 1997

Multiple comparisons theory and methods. Departament of statistics the Ohio State University. USA, 1996. Jason C. Hsu. Chapman Hall/CRC.

Practical Nonparametrics Statistics. W.J. Conover, 1999

Ramalho M.A.P., Ferreira D.F., Oliveira A.C. 2000. Experimentacao em Genetica e Melhoramento de Plantas. Editora UFLA.

Scott R.J., Knott M. 1974. A cluster analysis method for grouping mans in the analysis of variance. Biometrics, 30, 507-512.

See Also

```
DBC, DICT, DQLT
```

Examples

desc

Descriptive: Descriptive analysis

Description

Performs the descriptive analysis of an experiment with a factor of interest.

38 desc2fat

Usage

```
desc(trat, response, ylab = "Response", xlab = "Treatment", ylim = NA)
```

Arguments

trat Numerical or complex vector with treatments

response Numerical vector containing the response of the experiment.

ylab Variable response name (Accepts the *expression*() function)

xlab x name (Accepts the *expression*() function)

ylim y-axis scale

Value

The function returns exploratory measures of position and dispersion, such as mean, median, maximum, minimum, coefficient of variation, etc ...

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

See Also

```
desc2fat, tabledesc,dispvar
```

Examples

```
library(AgroR)
data("pomegranate")
with(pomegranate, desc(trat,WL))
```

desc2fat

Descriptive: Descriptive analysis (Two factors)

Description

It performs the descriptive analysis of an experiment with two factors of interest.

```
desc2fat(f1, f2, response, ylab = "Response", theme = theme_classic())
```

desc3fat 39

Arguments

f1	Numeric or complex vector with factor 1 levels
f2	Numeric or complex vector with factor 2 levels
	NT 1 1 4 4 1 1 C.1

response Numerical vector containing the response of the experiment. ylab Variable response name (Accepts the *expression*() function)

theme ggplot2 theme (default is theme_classic())

Value

The function returns exploratory measures of position and dispersion, such as mean, median, maximum, minimum, coefficient of variation, etc ...

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

Examples

```
library(AgroR)
data(cloro)
with(cloro, desc2fat(f1,f2,resp))
```

desc3fat

Descriptive: Descriptive analysis (Three factors)

Description

Performs the descriptive graphical analysis of an experiment with three factors of interest.

```
desc3fat(
  f1,
  f2,
  f3,
  response,
  legend.title = "Legend",
  xlab = "",
  ylab = "Response",
  theme = theme_classic(),
  plot = "interaction"
)
```

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Arguments

f1	Numeric or complex vector with factor 1 levels
f2	Numeric or complex vector with factor 2 levels
f3	Numeric or complex vector with factor 3 levels

response Numerical vector containing the response of the experiment.

legend.title Legend title

xlab x name (Accepts the *expression*() function)

ylab Variable response name (Accepts the *expression*() function)

theme ggplot theme

plot "interaction" or "box"

Value

The function returns a triple interaction graph.

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

Examples

```
library(AgroR)
data(enxofre)
with(enxofre, desc3fat(f1, f2, f3, resp))
```

desd_fat2_quant_ad Analysis: Regression analysis by orthogonal polynomials for double factorial scheme with additional control

Description

Regression analysis by orthogonal polynomials for double factorial scheme with additional control. Cases in which the additional belongs to the regression curve, being common to the qualitative levels. In these cases, the additional (usually dose 0/control treatment) is not part of the factor arrangement. One option addressed by this function is to analyze a priori as a double factorial scheme with an additional one and correct the information a posteriore using information from the initial analysis, such as the degree of freedom and the sum of squares of the residue.

```
desd_fat2_quant_ad(output, ad.value = 0, design = "FAT2DIC.ad", grau = 1)
```

desd_fat2_quant_ad 41

Arguments

output Output from a FAT2DIC.ad or FAT2DBC.ad function

ad.value Additional treatment quantitative factor level

design Type of experimental project (FAT2DIC.ad or FAT2DBC.ad)

grau Degree of the polynomial (only for the isolated effect of the quantitative factor)

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Leandro Simoes Azeredo Goncalves

Rodrigo Yudi Palhaci Marubayashi

Examples

```
# Data set
trat=rep(c("A", "B", "C"), e=12)
dose=rep(rep(c(200,400,600,800),e=3),3)
d0=c(40,45,48)
respo=c(60,55,56, 60,65,66, 70,75,76,
     80,85,86, 50,55,56, 70,75,76,
      60,65,66, 50,45,46, 50,45,46,
      50,55,66, 70,75,76, 80,85,86)
repe=rep(c("R1","R2","R3"),12)
# Analysis FAT2DIC.ad
resu=FAT2DIC.ad(trat,dose,repe=repe,respo,responseAd=d0,quali=c(TRUE,FALSE),grau21=c(1,2,1))
# Regression analysis
desd_fat2_quant_ad(resu,ad.value=0,design="FAT2DIC.ad")
# Data set
trat=rep(c("A", "B"), e=12)
dose=rep(rep(c(200,400,600,800),e=3),2)
d0=c(40,45,48)
respo=c(60,55,56,60,65,66,70,75,76,80,85,86,50,45,46,50,55,66,70,75,76,80,85,86)
repe=rep(c("R1","R2","R3"),8)
# Analysis FAT2DIC.ad
resu=FAT2DIC.ad(trat,dose,repe = repe,respo,responseAd = d0,quali = c(TRUE,FALSE))
# Regression analysis
desd_fat2_quant_ad(resu,ad.value=0,design="FAT2DIC.ad",grau=1)
```

DIC DIC

DIC

Analysis: Completely randomized design

Description

Statistical analysis of experiments conducted in a completely randomized and balanced design with a factor considering the fixed model. The function presents the option to use non-parametric method or transform the dataset.

```
DIC(
  trat,
  response,
  norm = "sw",
  homog = "bt",
  alpha.f = 0.05,
  alpha.t = 0.05,
  quali = TRUE,
 mcomp = "tukey",
  grau = 1,
  transf = 1,
  constant = 0,
  test = "parametric",
  mcompNP = "LSD",
  p.adj = "holm",
  geom = "bar",
  theme = theme_classic(),
  ylab = "Response",
  sup = NA,
  CV = TRUE,
  xlab = "",
  fill = "lightblue",
  angle = 0,
  family = "sans",
  textsize = 12,
  labelsize = 4,
  dec = 3,
  width.column = NULL,
  width.bar = 0.3,
  addmean = TRUE,
  errorbar = TRUE,
  posi = "top",
  point = "mean_sd",
  pointsize = 5,
  angle.label = 0,
  ylim = NA
```

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)

Arguments

trat Numerical or complex vector with treatments

response Numerical vector containing the response of the experiment.

norm Error normality test (default is Shapiro-Wilk)
homog Homogeneity test of variances (default is Bartlett)
alpha.f Level of significance of the F test (default is 0.05)

alpha.t Significance level of the multiple comparison test (*default* is 0.05)

quali Defines whether the factor is quantitative or qualitative (*default* is qualitative)
mcomp Multiple comparison test (Tukey (*default*), LSD, Scott-Knott and Duncan)

grau Degree of polynomial in case of quantitative factor (*default* is 1)

transf Applies data transformation (default is 1; for log consider 0, 'angular' for angu-

lar transformation)

constant Add a constant for transformation (enter value)

test "parametric" - Parametric test or "noparametric" - non-parametric test

mcompNP Multiple comparison test (LSD (default) or dunn)

p.adj Method for adjusting p values for Kruskal-Wallis ("none","holm","hommel",

"hochberg", "bonferroni", "BH", "BY", "fdr")

geom Graph type (columns, boxes or segments)
theme ggplot2 theme (default is theme classic())

ylab Variable response name (Accepts the *expression*() function)

Number of units above the standard deviation or average bar on the graph

CV Plotting the coefficient of variation and p-value of Anova (*default* is TRUE)

xlab Treatments name (Accepts the *expression*() function)

fill Defines chart color (to generate different colors for different treatments, define

fill = "trat")

angle x-axis scale text rotation

family Font family textsize Font size Label size

dec Number of cells

width.column Width column if geom="bar"

width.bar Width errorbar

addmean Plot the average value on the graph (*default* is TRUE)

errorbar Plot the standard deviation bar on the graph (In the case of a segment and column

graph) - default is TRUE

posi Legend position

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point Defines whether to plot mean ("mean"), mean with standard deviation ("mean_sd"

 $\hbox{-} \textit{default}) \ or \ mean \ with \ standard \ error \ ("mean_se"). \ For \ quali=FALSE \ or \ quali=TRUE.$

For parametric test it is possible to plot the square root of QMres (mean_qmres)

pointsize Point size angle.label label angle

ylim Define a numerical sequence referring to the y scale. You can use a vector or the

'seq' command.

Value

The table of analysis of variance, the test of normality of errors (Shapiro-Wilk ("sw"), Lilliefors ("li"), Anderson-Darling ("ad"), Cramer-von Mises ("cvm"), Pearson ("pearson") and Shapiro-Francia ("sf")), the test of homogeneity of variances (Bartlett ("bt") or Levene ("levene")), the test of independence of Durbin-Watson errors, the test of multiple comparisons (Tukey ("tukey"), LSD ("lsd"), Scott-Knott ("sk") or Duncan ("duncan")) or adjustment of regression models up to grade 3 polynomial, in the case of quantitative treatments. Non-parametric analysis can be used by the Kruskal-Wallis test. The column, segment or box chart for qualitative treatments is also returned. The function also returns a standardized residual plot.

Note

Enable ggplot2 package to change theme argument.

The ordering of the graph is according to the sequence in which the factor levels are arranged in the data sheet. The bars of the column and segment graphs are standard deviation.

Post hoc test in nonparametric is using the criterium Fisher's least significant difference (p-adj="holm").

CV and p-value of the graph indicate coefficient of variation and p-value of the F test of the analysis of variance.

In the final output when transformation (transf argument) is different from 1, the columns resp and respo in the mean test are returned, indicating transformed and non-transformed mean, respectively.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br> Leandro Simoes Azeredo Goncalves

Rodrigo Yudi Palhaci Marubayashi

References

Principles and procedures of statistics a biometrical approach Steel, Torry and Dickey. Third Edition 1997

Multiple comparisons theory and methods. Departament of statistics the Ohio State University. USA, 1996. Jason C. Hsu. Chapman Hall/CRC.

W.J. Conover, Practical Nonparametrics Statistics. 1999

Ramalho M.A.P., Ferreira D.F., Oliveira A.C. 2000. Experimentacao em Genetica e Melhoramento de Plantas. Editora UFLA.

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Scott R.J., Knott M. 1974. A cluster analysis method for grouping mans in the analysis of variance. Biometrics, 30, 507-512.

Mendiburu, F., and de Mendiburu, M. F. (2019). Package 'agricolae'. R Package, Version, 1-2.

Hothorn, T. et al. Package 'lmtest'. Testing linear regression models. https://cran. r-project. org/web/packages/lmtest/lmtest. pdf. Accessed, v. 6, 2015.

See Also

DBC DQL

Examples

```
library(AgroR)
data(pomegranate)
with(pomegranate, DIC(trat, WL, ylab = "Weight loss (%)")) # tukey
with(pomegranate, DIC(trat, WL, mcomp = "sk", ylab = "Weight loss (%)"))
with(pomegranate, DIC(trat, WL, mcomp = "duncan", ylab = "Weight loss (%)"))
# Kruskal-Wallis
with(pomegranate, DIC(trat, WL, test = "noparametric", ylab = "Weight loss (%)"))
# chart type
with(pomegranate, DIC(trat, WL, geom="point", ylab = "Weight loss (%)"))
with(pomegranate, DIC(trat, WL, ylab = "Weight loss (%)", xlab="Treatments"))
# quantitative factor
data("phao")
with(phao, DIC(dose,comp,quali=FALSE,grau=2,
           xlab = expression("Dose"~(g~vase^-1)),
           ylab="Leaf length (cm)"))
# data transformation
#===========
data("pepper")
with(pepper, DIC(Acesso, VitC, transf = 0,ylab="Vitamin C"))
```

Analysis: Completely randomized design with an additional treatment for quantitative factor

46 dic.ad

Description

Statistical analysis of experiments conducted in a completely randomized with an additional treatment and balanced design with a factor considering the fixed model.

Usage

```
dic.ad(
  trat,
  response,
  responsead,
  grau = 1,
  norm = "sw",
  homog = "bt",
  alpha.f = 0.05,
  theme = theme_classic(),
  ylab = "response",
  xlab = "independent",
  family = "sans",
  posi = "top",
  pointsize = 4.5,
  linesize = 0.8,
  width.bar = NA,
  point = "mean_sd"
)
```

Arguments

trat	Numerical or complex vector with treatments
response	Numerical vector containing the response of the experiment.
responsead	Numerical vector with additional treatment responses
grau	Degree of polynomial in case of quantitative factor (default is 1)
norm	Error normality test (default is Shapiro-Wilk)
homog	Homogeneity test of variances (default is Bartlett)
alpha.f	Level of significance of the F test (<i>default</i> is 0.05)
theme	ggplot2 theme (default is theme_classic())
ylab	Variable response name (Accepts the expression() function)
xlab	Treatments name (Accepts the expression() function)
family	Font family
posi	Legend position
pointsize	Point size
linesize	line size (Trendline and Error Bar)
width.bar	width of the error bars of a regression graph.
point	Defines whether to plot mean ("mean"), mean with standard deviation ("mean_sd" - <i>default</i>) or mean with standard error (<i>default</i> - "mean_se"). For quali=FALSE or quali=TRUE.

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Value

The table of analysis of variance, the test of normality of errors (Shapiro-Wilk ("sw"), Lilliefors ("li"), Anderson-Darling ("ad"), Cramer-von Mises ("cvm"), Pearson ("pearson") and Shapiro-Francia ("sf")), the test of homogeneity of variances (Bartlett ("bt") or Levene ("levene")), the test of independence of Durbin-Watson errors, adjustment of regression models up to grade 3 polynomial. The function also returns a standardized residual plot.

Note

In some experiments, the researcher may study a quantitative factor, such as fertilizer doses, and present a control, such as a reference fertilizer, treated as a qualitative control. In these cases, there is a difference between considering only the residue in the unfolding of the polynomial, removing or not the qualitative treatment, or since a treatment is excluded from the analysis. In this approach, the residue used is also considering the qualitative treatment, a method similar to the factorial scheme with additional control.

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

Examples

```
datadicad=data.frame(doses = c(rep(c(1:5),e=3)),

resp = c(3,4,3,5,5,6,7,7,8,4,4,5,2,2,3))

with(datadicad,dic.ad(doses, resp, rnorm(3,6,0.1),grau=2))
```

DIC.glm

Analysis: Completely randomized design by glm

Description

Statistical analysis of experiments conducted in a completely randomized design using a generalized linear model. It performs the deviance analysis and the effect is tested by a chi-square test. Multiple comparisons are adjusted by Tukey.

```
DIC.glm(
  trat,
  response,
  glm.family = "binomial",
  quali = TRUE,
  alpha.f = 0.05,
  alpha.t = 0.05,
  geom = "bar",
```

DIC.glm

```
theme = theme_classic(),
 sup = NA,
 ylab = "Response",
 xlab = "",
 fill = "lightblue",
 angle = 0,
 family = "sans",
 textsize = 12,
 labelsize = 5,
 dec = 3,
 addmean = TRUE,
 errorbar = TRUE,
 posi = "top",
 point = "mean_sd",
 angle.label = 0
)
```

Arguments

trat	Numerical or complex vector with treatments
response	Numerical vector containing the response of the experiment. Use cbind(resp, n-resp) for binomial or quasibinomial family.
glm.family	distribution family considered (default is binomial)
quali	Defines whether the factor is quantitative or qualitative (<i>default</i> is qualitative)
alpha.f	Level of significance of the F test (<i>default</i> is 0.05)
alpha.t	Significance level of the multiple comparison test (<i>default</i> is 0.05)
geom	Graph type (columns, boxes or segments)
theme	ggplot2 theme (default is theme_classic())
sup	Number of units above the standard deviation or average bar on the graph
ylab	Variable response name (Accepts the expression() function)
xlab	Treatments name (Accepts the expression() function)
fill	Defines chart color (to generate different colors for different treatments, define fill = "trat")
angle	x-axis scale text rotation
family	Font family
textsize	Font size
labelsize	Label size
dec	Number of cells
addmean	Plot the average value on the graph (default is TRUE)
errorbar	Plot the standard deviation bar on the graph (In the case of a segment and column graph) - <i>default</i> is TRUE
posi	Legend position
point	Defines whether to plot mean ("mean"), mean with standard deviation ("mean_sd" - <i>default</i>) or mean with standard error (<i>default</i> - "mean_se").
angle.label	label angle

DICT 49

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br> Leandro Simoes Azeredo Goncalves Rodrigo Yudi Palhaci Marubayashi

Examples

DICT

Analysis: Completely randomized design evaluated over time

Description

Function of the AgroR package for the analysis of experiments conducted in a completely randomized, qualitative, uniform qualitative design with multiple assessments over time, however without considering time as a factor.

```
DICT(
   trat,
   time,
   response,
   alpha.f = 0.05,
   alpha.t = 0.05,
   mcomp = "tukey",
   theme = theme_classic(),
   geom = "bar",
```

DICT

```
xlab = "Independent",
 ylab = "Response",
 p.adj = "holm",
 dec = 3,
 fill = "gray",
 error = TRUE,
 textsize = 12,
 labelsize = 5,
  pointsize = 4.5,
  family = "sans",
  sup = 0,
  addmean = FALSE,
 legend = "Legend",
 ylim = NA,
 width.bar = 0.2,
 size.bar = 0.8,
 posi = c(0.1, 0.8),
 xnumeric = FALSE,
 all.letters = FALSE
)
```

Arguments

trat	Numerical or complex vector with treatments
time	Numerical or complex vector with times
response	Numerical vector containing the response of the experiment.
alpha.f	Level of significance of the F test (default is 0.05)
alpha.t	Significance level of the multiple comparison test (default is 0.05)
mcomp	Multiple comparison test (Tukey (<i>default</i>), LSD ("lsd"), Scott-Knott ("sk"), Duncan ("duncan") and Kruskal-Wallis ("kw"))
theme	ggplot2 theme (default is theme_classic())
geom	Graph type (columns - "bar" or segments "point")
xlab	treatments name (Accepts the expression() function)
ylab	Variable response name (Accepts the expression() function)
p.adj	Method for adjusting p values for Kruskal-Wallis ("none", "holm", "hommel", "hochberg", "bonferroni", "BH", "BY", "fdr")
dec	Number of cells
fill	Defines chart color (to generate different colors for different treatments, define fill = "trat")
error	Add error bar
textsize	Font size of the texts and titles of the axes
labelsize	Font size of the labels
pointsize	Point size
family	Font family

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sup Number of units above the standard deviation or average bar on the graph

addmean Plot the average value on the graph (*default* is TRUE)

legend Legend title

ylim Define a numerical sequence referring to the y scale. You can use a vector or the

'seq' command.

width.bar width error bar size.bar size error bar posi Legend position

xnumeric Declare x as numeric (*default* is FALSE)

all.letters Adds all label letters regardless of whether it is significant or not.

Value

The function returns the p-value of Anova, the assumptions of normality of errors, homogeneity of variances and independence of errors, multiple comparison test, as well as a line graph

Note

The ordering of the graph is according to the sequence in which the factor levels are arranged in the data sheet. The bars of the column and segment graphs are standard deviation.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Leandro Simoes Azeredo Goncalves

Rodrigo Yudi Palhaci Marubayashi

References

Principles and procedures of statistics a biometrical approach Steel, Torry and Dickey. Third Edition 1997

Multiple comparisons theory and methods. Departament of statistics the Ohio State University. USA, 1996. Jason C. Hsu. Chapman Hall/CRC.

Practical Nonparametrics Statistics. W.J. Conover, 1999

Ramalho M.A.P., Ferreira D.F., Oliveira A.C. 2000. Experimentacao em Genetica e Melhoramento de Plantas. Editora UFLA.

Scott R.J., Knott M. 1974. A cluster analysis method for grouping mans in the analysis of variance. Biometrics, 30, 507-512.

See Also

DIC, DBCT, DQLT

52 dispvar

Examples

```
rm(list=ls())
data(simulate1)
attach(simulate1)
with(simulate1, DICT(trat, tempo, resp))
with(simulate1, DICT(trat, tempo, resp, fill="rainbow",family="serif"))
with(simulate1, DICT(trat, tempo, resp,geom="bar",sup=40))
with(simulate1, DICT(trat, tempo, resp,geom="point",sup=40))
```

dispvar

Descriptive: Boxplot with standardized data

Description

It makes a graph with the variables and/or treatments with the standardized data.

Usage

```
dispvar(
  data,
  trat = NULL,
  theme = theme_bw(),
  ylab = "Standard mean",
  xlab = "Variable",
  family = "serif",
  textsize = 12,
  fill = "lightblue"
)
```

Arguments

data	data.frame containing the response of the experiment.
trat	Numerical or complex vector with treatments
theme	ggplot2 theme (default is theme_bw())
ylab	Variable response name (Accepts the expression() function)
xlab	Treatments name (Accepts the expression() function)
family	Font family
textsize	Font size
fill	Defines chart color

Value

Returns a chart of boxes with standardized data

DQL 53

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

Examples

```
library(AgroR)
data("pomegranate")
dispvar(pomegranate[,-1])
trat=pomegranate$trat
dispvar(pomegranate[,-1], trat)
```

DQL

Analysis: Latin square design

Description

This is a function of the AgroR package for statistical analysis of experiments conducted in Latin Square and balanced design with a factor considering the fixed model.

```
DQL(
  trat,
  line,
  column,
  response,
  norm = "sw",
  homog = "bt",
  alpha.f = 0.05,
  alpha.t = 0.05,
  quali = TRUE,
 mcomp = "tukey",
  grau = 1,
  transf = 1,
  constant = 0,
  geom = "bar",
  theme = theme_classic(),
  sup = NA,
  CV = TRUE,
  ylab = "Response",
  xlab = "",
  textsize = 12,
  labelsize = 4,
  fill = "lightblue",
```

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```
angle = 0,
family = "sans",
dec = 3,
width.column = NULL,
width.bar = 0.3,
addmean = TRUE,
errorbar = TRUE,
posi = "top",
point = "mean_sd",
pointsize = 5,
angle.label = 0,
ylim = NA
```

Arguments

trat Numerical or complex vector with treatments
line Numerical or complex vector with lines
column Numerical or complex vector with columns

response Numerical vector containing the response of the experiment.

norm Error normality test (default is Shapiro-Wilk)
homog Homogeneity test of variances (default is Bartlett)
alpha.f Level of significance of the F test (default is 0.05)

alpha.t Significance level of the multiple comparison test (*default* is 0.05)

quali Defines whether the factor is quantitative or qualitative (*default* is qualitative)
mcomp Multiple comparison test (Tukey (*default*), LSD, Scott-Knott and Duncan)

grau Degree of polynomial in case of quantitative factor (*default* is 1)

transf Applies data transformation (default is 1; for log consider 0; 'angular' for angu-

lar transformation)

constant Add a constant for transformation (enter value)
geom Graph type (columns, boxes or segments)
theme ggplot2 theme (default is theme_classic())

Number of units above the standard deviation or average bar on the graph

CV Plotting the coefficient of variation and p-value of Anova (*default* is TRUE)

ylab Variable response name (Accepts the *expression*() function)
xlab Treatments name (Accepts the *expression*() function)

textsize Font size labelsize Label size

fill Defines chart color (to generate different colors for different treatments, define

fill = "trat")

angle x-axis scale text rotation

family Font family

dec Number of cells

width.column Width column if geom="bar"

width.bar Width errorbar

addmean Plot the average value on the graph (*default* is TRUE)

errorbar Plot the standard deviation bar on the graph (In the case of a segment and column

graph) - default is TRUE

posi Legend position

point Defines whether to plot mean ("mean"), mean with standard deviation ("mean sd"

- default) or mean with standard error ("mean_se"). For parametric test it is pos-

sible to plot the square root of QMres (mean_qmres).

pointsize Point size angle.label label angle

ylim Define a numerical sequence referring to the y scale. You can use a vector or the

'seq' command.

Value

The table of analysis of variance, the test of normality of errors (Shapiro-Wilk ("sw"), Lilliefors ("li"), Anderson-Darling ("ad"), Cramer-von Mises ("cvm"), Pearson ("pearson") and Shapiro-Francia ("sf")), the test of homogeneity of variances (Bartlett ("bt") or Levene ("levene")), the test of independence of Durbin-Watson errors, the test of multiple comparisons (Tukey ("tukey"), LSD ("lsd"), Scott-Knott ("sk") or Duncan ("duncan")) or adjustment of regression models up to grade 3 polynomial, in the case of quantitative treatments. The column, segment or box chart for qualitative treatments is also returned. The function also returns a standardized residual plot.

Note

The ordering of the graph is according to the sequence in which the factor levels are arranged in the data sheet. The bars of the column and segment graphs are standard deviation.

CV and p-value of the graph indicate coefficient of variation and p-value of the F test of the analysis of variance.

In the final output when transformation (transf argument) is different from 1, the columns resp and respo in the mean test are returned, indicating transformed and non-transformed mean, respectively.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Leandro Simoes Azeredo Goncalves

Rodrigo Yudi Palhaci Marubayashi

References

Principles and procedures of statistics a biometrical approach Steel, Torry and Dickey. Third Edition 1997

DQLT

Multiple comparisons theory and methods. Departament of statistics the Ohio State University. USA, 1996. Jason C. Hsu. Chapman Hall/CRC.

Ramalho M.A.P., Ferreira D.F., Oliveira A.C. 2000. Experimentacao em Genetica e Melhoramento de Plantas. Editora UFLA.

Scott R.J., Knott M. 1974. A cluster analysis method for grouping mans in the analysis of variance. Biometrics, 30, 507-512.

Mendiburu, F., and de Mendiburu, M. F. (2019). Package 'agricolae'. R Package, Version, 1-2.

See Also

```
DIC, DBC
```

Examples

```
library(AgroR)
data(porco)
with(porco, DQL(trat, linhas, colunas, resp, ylab="Weigth (kg)"))
```

DQLT

Analysis: Latin square design evaluated over time

Description

Function of the AgroR package for the analysis of experiments conducted in a balanced qualitative single-square Latin design with multiple assessments over time, however without considering time as a factor.

```
DQLT(
  trat.
  line,
  column,
  time,
  response,
  alpha.f = 0.05,
  alpha.t = 0.05,
  mcomp = "tukey",
  error = TRUE,
  xlab = "Independent",
  ylab = "Response",
  textsize = 12,
  labelsize = 5,
  pointsize = 4.5,
  family = "sans",
  sup = 0,
  addmean = FALSE,
```

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```
posi = c(0.1, 0.8),
  geom = "bar",
  fill = "gray",
  legend = "Legend",
  ylim = NA,
 width.bar = 0.2,
  size.bar = 0.8,
  dec = 3,
  theme = theme_classic(),
  xnumeric = FALSE,
  all.letters = FALSE
)
```

Arguments

trat

Numerical or complex vector with treatments line Numerical or complex vector with line column Numerical or complex vector with column time Numerical or complex vector with times Numerical vector containing the response of the experiment. response alpha.f Level of significance of the F test (*default* is 0.05) Significance level of the multiple comparison test (*default* is 0.05) alpha.t Multiple comparison test (Tukey (default), LSD, Scott-Knott and Duncan) mcomp Add error bar (SD) error xlab Treatments name (Accepts the *expression*() function) ylab Variable response name (Accepts the *expression*() function) Font size of the texts and titles of the axes textsize labelsize Font size of the labels pointsize Point size family Font family sup Number of units above the standard deviation or average bar on the graph addmean Plot the average value on the graph (*default* is TRUE) posi Legend position Graph type (columns - "bar" or segments "point") geom fill Defines chart color (to generate different colors for different treatments, define fill = "trat") legend Legend title ylim Define a numerical sequence referring to the y scale. You can use a vector or the 'seq' command. width.bar width error bar size.bar size error bar

58 DQLT

dec Number of cells

theme ggplot2 theme (default is theme_classic())
xnumeric Declare x as numeric (default is FALSE)

all.letters Adds all label letters regardless of whether it is significant or not.

Details

The p-value of the analysis of variance, the normality test for Shapiro-Wilk errors, the Bartlett homogeneity test of variances, the independence of Durbin-Watson errors and the multiple comparison test (Tukey, Scott-Knott, LSD or Duncan).

Value

The function returns the p-value of Anova, the assumptions of normality of errors, homogeneity of variances and independence of errors, multiple comparison test, as well as a line graph

Note

The ordering of the graph is according to the sequence in which the factor levels are arranged in the data sheet. The bars of the column and segment graphs are standard deviation.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Leandro Simoes Azeredo Goncalves

Rodrigo Yudi Palhaci Marubayashi

References

Principles and procedures of statistics a biometrical approach Steel, Torry and Dickey. Third Edition 1997

Multiple comparisons theory and methods. Departament of statistics the Ohio State University. USA, 1996. Jason C. Hsu. Chapman Hall/CRC.

Practical Nonparametrics Statistics. W.J. Conover, 1999

Ramalho M.A.P., Ferreira D.F., Oliveira A.C. 2000. Experimentacao em Genetica e Melhoramento de Plantas. Editora UFLA.

Scott R.J., Knott M. 1974. A cluster analysis method for grouping mans in the analysis of variance. Biometrics, 30, 507-512.

See Also

DQL, DICT, DBCT

dunn 59

Examples

```
rm(list=ls())
data(simulate3)
attach(simulate3)
DQLT(trat, linhas, colunas, tempo, resp)
```

dunn

Analysis: Post-hoc Dunn

Description

Perform Kruskal wallis and dunn post-hoc test

Usage

```
dunn(trat, resp, method = "holm", alpha = 0.05, decreasing = TRUE)
```

Arguments

trat Numerical or complex vector with treatments

resp Vector with response

method the p-value for multiple comparisons ("none", "bonferroni", "sidak", "holm",

"hs", "hochberg", "bh", "by"). The default is no adjustment for multiple com-

parisons

alpha Significance level of the post-hoc (*default* is 0.05)

decreasing Should the order of the letters be increasing or decreasing.

Value

Kruskal-wallis and dunn's post-hoc test returns

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Examples

```
library(AgroR)
data(pomegranate)
with(pomegranate, dunn(trat, WL))
```

dunnett

dunnett

Analysis: Dunnett test

Description

The function performs the Dunnett test

Usage

```
dunnett(
  trat,
  resp,
  control,
  model = "DIC",
  block = NA,
  column = NA,
  line = NA,
  alpha.t = 0.05,
  pointsize = 5,
  pointshape = 21,
  linesize = 1,
  labelsize = 4,
  textsize = 12,
  errorsize = 1,
  widthsize = 0.2,
  label = "Response",
  fontfamily = "sans"
)
```

Arguments

trat	Numerical or complex vector with treatments
resp	Numerical vector containing the response of the experiment.
control	Treatment considered control (write identical to the name in the vector)
model	Experimental design (DIC, DBC or DQL)
block	Numerical or complex vector with blocks
column	Numerical or complex vector with columns
line	Numerical or complex vector with lines
alpha.t	Significance level (default is 0.05)
pointsize	Point size
pointshape	Shape
linesize	Line size
labelsize	Label size

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textsize	Font size
errorsize	Errorbar size
widthsize	Width errorbar
label	Variable label
fontfamily	font family

Value

I return the Dunnett test for experiments in a completely randomized design, randomized blocks or Latin square.

Note

Do not use the "-" symbol or space in treatment names

Examples

```
# complete randomized design
data("pomegranate")
with(pomegranate,dunnett(trat=trat,resp=WL,control="T1"))
# randomized block design in factorial double
library(AgroR)
data(cloro)
attach(cloro)
respAd=c(268, 322, 275, 350, 320)
a=FAT2DBC.ad(f1, f2, bloco, resp, respAd,
         ylab="Number of nodules",
         legend = "Stages", mcomp="sk")
data=rbind(data.frame(trat=paste(f1,f2,sep = ""),bloco=bloco,resp=resp),
       data.frame(trat=c("Test","Test","Test","Test"),
               bloco=unique(bloco),resp=respAd))
with(data,dunnett(trat = trat,
             resp = resp,
             control = "Test",
             block=bloco,model = "DBC"))
```

emerg

Dataset: Emergence of passion fruit seeds over time.

Description

The data come from an experiment conducted at the State University of Londrina, aiming to study the emergence of yellow passion fruit seeds over time. Data are partial from one of the treatments studied. Four replicates with eight seeds each were used.

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Usage

```
data("emerg")
```

Format

data.frame containing data set

time numeric vector with time

resp Numeric vector with emergence

See Also

aristolochia, cloro, laranja, enxofre, laranja, mirtilo, passiflora, phao, porco, pomegranate, simulate1, simulate2, simulate3, tomate, weather

Examples

data(emerg)

enxofre

Dataset: Sulfur data

Description

The experiment was carried out in a randomized block design in a $3 \times 3 \times 3$ triple factorial scheme: syrup volume (75, 225 and 675 L), sulfur doses (150, 450, 1350) and time of application (vegetative, complete cycle and reproductive system) with four repetitions. Yield in kg / ha of soybean was evaluated.

Usage

```
data(enxofre)
```

Format

data.frame containing data set

- f1 Categorical vector with factor 1
- f2 Categorical vector with factor 2
- f2 Categorical vector with factor 3

bloco Categorical vector with block

resp Numeric vector

See Also

cloro, laranja, mirtilo, pomegranate, porco, sensorial, simulate1, simulate2, simulate3, tomate, weather, phao, passiflora, aristolochia

eucalyptus 63

Examples

```
data(enxofre)
```

eucalyptus

Dataset: Eucaliptus grandis Barbin (2013)

Description

The data refer to the height in meters of *Eucalyptus grandis* plants, with 7 years of age, from three trials (Araraquara - Exp 1; Bento Quintino - Exp 2; Mogi-Guacu - Exp 3) in randomized blocks, under 6 progenies. The data were taken from the book by Decio Barbin (2013) and are from the Instituto Florestal de Tupi/SP.

Usage

```
data("eucalyptus")
```

Format

data.frame containing data set

trati Categorical vector with treatments

bloc Categorical vector with block

exp Categorical vector with experiment

resp Numeric vector

References

Planejamento e Analise Estatistica de Experimentos Agronomicos (2013) - Decio Barbin - pg. 177

See Also

cloro, enxofre, laranja, pomegranate, porco, sensorial, simulate1, simulate2, simulate3, tomate, weather

Examples

```
data(eucalyptus)
```

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FAT2DBC

Analysis: DBC experiments in double factorial

Description

Analysis of an experiment conducted in a randomized block design in a double factorial scheme using analysis of variance of fixed effects.

```
FAT2DBC(
  f1,
  f2,
  block,
  response,
  norm = "sw",
  homog = "bt",
  alpha.f = 0.05,
  alpha.t = 0.05,
  quali = c(TRUE, TRUE),
  names.fat = c("F1", "F2"),
  mcomp = "tukey",
  grau = c(NA, NA),
  grau12 = NA,
  grau21 = NA,
  transf = 1,
  constant = 0,
  geom = "bar",
  theme = theme_classic(),
  ylab = "Response",
  xlab = "",
  xlab.factor = c("F1", "F2"),
  legend = "Legend",
  fill = "lightblue",
  angle = 0,
  textsize = 12,
  labelsize = 4,
  dec = 3,
  width.column = 0.9,
  width.bar = 0.3,
  family = "sans",
  point = "mean_sd",
  addmean = TRUE,
  errorbar = TRUE,
  CV = TRUE,
  sup = NA,
  color = "rainbow",
```

FAT2DBC 65

```
posi = "right",
ylim = NA,
angle.label = 0
)
```

Arguments

Numeric or complex vector with factor 1 levels
 Numeric or complex vector with factor 2 levels
 Numerical or complex vector with blocks

response Numerical vector containing the response of the experiment.

norm Error normality test (default is Shapiro-Wilk)
homog Homogeneity test of variances (default is Bartlett)
alpha.f Level of significance of the F test (default is 0.05)

alpha.t Significance level of the multiple comparison test (*default* is 0.05) quali Defines whether the factor is quantitative or qualitative (*qualitative*)

names.fat Name of factors

mcomp Multiple comparison test (Tukey (*default*), LSD, Scott-Knott and Duncan)

grau Polynomial degree in case of quantitative factor (*default* is 1). Provide a vector

with two elements.

grau12 Polynomial degree in case of quantitative factor (*default* is 1). Provide a vector

with n levels of factor 2, in the case of interaction f1 x f2 and qualitative factor

2 and quantitative factor 1.

grau21 Polynomial degree in case of quantitative factor (*default* is 1). Provide a vector

with n levels of factor 1, in the case of interaction f1 x f2 and qualitative factor

1 and quantitative factor 2.

transf Applies data transformation (default is 1; for log consider 0; 'angular' for angu-

lar transformation)

constant Add a constant for transformation (enter value)

geom Graph type (columns or segments (For simple effect only))

theme ggplot2 theme (default is theme_classic())

ylab Variable response name (Accepts the *expression*() function)

xlab Treatments name (Accepts the *expression*() function)

xlab. factor Provide a vector with two observations referring to the x-axis name of factors 1

and 2, respectively, when there is an isolated effect of the factors. This argument

uses 'parse'.

legend Legend title name

fill Defines chart color (to generate different colors for different treatments, define

fill = "trat")

angle x-axis scale text rotation

textsize font size

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labelsize label size

dec number of cells

width.column Width column if geom="bar"

width.bar Width errorbar family font family

point This function defines whether the point must have all points ("all"), mean ("mean"),

standard deviation (*default* - "mean_sd") or mean with standard error ("mean_se") if quali= FALSE. For quali=TRUE, 'mean_sd' and 'mean_se' change which in-

formation will be displayed in the error bar.

addmean Plot the average value on the graph (*default* is TRUE)

errorbar Plot the standard deviation bar on the graph (In the case of a segment and column

graph) - default is TRUE

CV Plotting the coefficient of variation and p-value of Anova (*default* is TRUE)

sup Number of units above the standard deviation or average bar on the graph

color Column chart color (*default* is "rainbow")

posi Legend position
ylim y-axis scale
angle.label label angle

Value

The table of analysis of variance, the test of normality of errors (Shapiro-Wilk, Lilliefors, Anderson-Darling, Cramer-von Mises, Pearson and Shapiro-Francia), the test of homogeneity of variances (Bartlett or Levene), the test of independence of Durbin-Watson errors, the test of multiple comparisons (Tukey, LSD, Scott-Knott or Duncan) or adjustment of regression models up to grade 3 polynomial, in the case of quantitative treatments. The column chart for qualitative treatments is also returned.

Note

The order of the chart follows the alphabetical pattern. Please use 'scale_x_discrete' from package ggplot2, 'limits' argument to reorder x-axis. The bars of the column and segment graphs are standard deviation.

The function does not perform multiple regression in the case of two quantitative factors.

In the final output when transformation (transf argument) is different from 1, the columns resp and respo in the mean test are returned, indicating transformed and non-transformed mean, respectively.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Leandro Simoes Azeredo Goncalves

Rodrigo Yudi Palhaci Marubayashi

References

Principles and procedures of statistics a biometrical approach Steel, Torry and Dickey. Third Edition 1997

Multiple comparisons theory and methods. Departament of statistics the Ohio State University. USA, 1996. Jason C. Hsu. Chapman Hall/CRC.

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Ramalho M.A.P., Ferreira D.F., Oliveira A.C. 2000. Experimentacao em Genetica e Melhoramento de Plantas. Editora UFLA.

Scott R.J., Knott M. 1974. A cluster analysis method for grouping mans in the analysis of variance. Biometrics, 30, 507-512.

Mendiburu, F., and de Mendiburu, M. F. (2019). Package 'agricolae'. R Package, Version, 1-2.

See Also

FAT2DBC.ad

Examples

```
# Example cloro
library(AgroR)
data(cloro)
attach(cloro)
FAT2DBC(f1, f2, bloco, resp, ylab="Number of nodules", legend = "Stages")
FAT2DBC(f1, f2, bloco, resp, mcomp="sk", ylab="Number of nodules", legend = "Stages")
# Example covercrops
library(AgroR)
data(covercrops)
attach(covercrops)
FAT2DBC(A, B, Bloco, Resp, ylab=expression("Yield"~(Kg~"100 m"^2)),
legend = "Cover crops")
FAT2DBC(A, B, Bloco, Resp, mcomp="sk", ylab=expression("Yield"~(Kg~"100 m"^2)),
legend = "Cover crops")
```

FAT2DBC.ad

Analysis: DBC experiment in double factorial design with an additional treatment

Description

Analysis of an experiment conducted in a randomized block design in a double factorial scheme using analysis of variance of fixed effects.

Usage

```
FAT2DBC.ad(
  f1,
  f2,
  block,
  response,
  responseAd,
  norm = "sw",
  homog = "bt",
  alpha.f = 0.05,
  alpha.t = 0.05,
  quali = c(TRUE, TRUE),
  names.fat = c("F1", "F2"),
 mcomp = "tukey",
  grau = c(NA, NA),
  grau12 = NA,
 grau21 = NA,
  transf = 1,
  constant = 0,
  geom = "bar",
  theme = theme_classic(),
  ylab = "Response",
  xlab = "",
  xlab.factor = c("F1", "F2"),
  legend = "Legend",
  ad.label = "Additional",
  color = "rainbow",
  fill = "lightblue",
  textsize = 12,
  labelsize = 4,
  addmean = TRUE,
  errorbar = TRUE,
  CV = TRUE,
  dec = 3,
 width.column = 0.9,
 width.bar = 0.3,
  angle = 0,
  posi = "right",
  family = "sans",
  point = "mean_sd",
  sup = NA,
 ylim = NA,
  angle.label = 0
)
```

Arguments

Numeric or complex vector with factor 1 levels

f2 Numeric or complex vector with factor 2 levels block Numeric or complex vector with repetitions

response Numerical vector containing the response of the experiment.

responseAd Numerical vector with additional treatment responses

norm Error normality test (default is Shapiro-Wilk)
homog Homogeneity test of variances (default is Bartlett)
alpha.f Level of significance of the F test (default is 0.05)

alpha.t Significance level of the multiple comparison test (*default* is 0.05) quali Defines whether the factor is quantitative or qualitative (*qualitative*)

names.fat Name of factors

mcomp Multiple comparison test (Tukey (*default*), LSD and Duncan)

grau Polynomial degree in case of quantitative factor (*default* is 1). Provide a vector

with two elements.

grau12 Polynomial degree in case of quantitative factor (*default* is 1). Provide a vector

with n levels of factor 2, in the case of interaction f1 x f2 and qualitative factor

2 and quantitative factor 1.

grau21 Polynomial degree in case of quantitative factor (*default* is 1). Provide a vector

with n levels of factor 1, in the case of interaction f1 x f2 and qualitative factor

1 and quantitative factor 2.

transf Applies data transformation (default is 1; for log consider 0; 'angular' for angu-

lar transformation)

constant Add a constant for transformation (enter value)

geom Graph type (columns or segments (For simple effect only))

theme ggplot2 theme (*default* is theme_classic())

ylab Variable response name (Accepts the *expression*() function)

xlab Treatments name (Accepts the *expression*() function)

xlab.factor Provide a vector with two observations referring to the x-axis name of factors 1

and 2, respectively, when there is an isolated effect of the factors. This argument

uses 'parse'.

legend Legend title name ad.label Aditional label

color Column chart color (*default* is "rainbow")

fill Defines chart color (to generate different colors for different treatments, define

fill = "trat")

textsize Font size labelsize Label Size

addmean Plot the average value on the graph (*default* is TRUE)

errorbar Plot the standard deviation bar on the graph (In the case of a segment and column

graph) - default is TRUE

CV Plotting the coefficient of variation and p-value of Anova (*default* is TRUE)

dec Number of cells

width.column Width column if geom="bar"

width.bar Width errorbar

angle x-axis scale text rotation

posi legend position family Font family

point This function defines whether the point must have all points ("all"), mean ("mean"),

standard deviation (*default* - "mean_sd") or mean with standard error ("mean_se") if quali= FALSE. For quali=TRUE, 'mean_sd' and 'mean_se' change which in-

formation will be displayed in the error bar.

sup Number of units above the standard deviation or average bar on the graph

ylim y-axis scale angle.label label angle

Value

The table of analysis of variance, the test of normality of errors (Shapiro-Wilk, Lilliefors, Anderson-Darling, Cramer-von Mises, Pearson and Shapiro-Francia), the test of homogeneity of variances (Bartlett or Levene), the test of independence of Durbin-Watson errors, the test of multiple comparisons (Tukey, LSD, Scott-Knott or Duncan) or adjustment of regression models up to grade 3 polynomial, in the case of quantitative treatments. The column chart for qualitative treatments is also returned.

Note

The order of the chart follows the alphabetical pattern. Please use 'scale_x_discrete' from package ggplot2, 'limits' argument to reorder x-axis. The bars of the column and segment graphs are standard deviation.

The function does not perform multiple regression in the case of two quantitative factors.

The assumptions of variance analysis disregard additional treatment

In the final output when transformation (transf argument) is different from 1, the columns resp and respo in the mean test are returned, indicating transformed and non-transformed mean, respectively.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br> Leandro Simoes Azeredo Goncalves

Rodrigo Yudi Palhaci Marubayashi

References

Principles and procedures of statistics a biometrical approach Steel, Torry and Dickey. Third Edition 1997

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Mendiburu, F., and de Mendiburu, M. F. (2019). Package 'agricolae'. R Package, Version, 1-2.

See Also

FAT2DBC

dunnett

Examples

```
library(AgroR)
data(cloro)
respAd=c(268, 322, 275, 350, 320)
with(cloro, FAT2DBC.ad(f1, f2, bloco, resp, respAd, ylab="Number of nodules", legend = "Stages"))
```

FAT2DIC

Analysis: DIC experiments in double factorial

Description

Analysis of an experiment conducted in a completely randomized design in a double factorial scheme using analysis of variance of fixed effects.

```
FAT2DIC(
  f1,
  f2,
  response,
  norm = "sw"
  homog = "bt",
  alpha.f = 0.05,
  alpha.t = 0.05,
  quali = c(TRUE, TRUE),
  names.fat = c("F1", "F2"),
  mcomp = "tukey",
  grau = c(NA, NA),
  grau12 = NA,
  grau21 = NA,
  transf = 1,
  constant = 0,
  geom = "bar",
```

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```
theme = theme_classic(),
  ylab = "Response",
  xlab = "",
  xlab.factor = c("F1", "F2"),
  legend = "Legend",
  color = "rainbow",
  fill = "lightblue",
  textsize = 12,
 labelsize = 4,
 addmean = TRUE,
 errorbar = TRUE,
 CV = TRUE,
 dec = 3,
 width.column = 0.9,
 width.bar = 0.3,
  angle = 0,
 posi = "right",
  family = "sans",
 point = "mean_sd",
  sup = NA,
 ylim = NA,
 angle.label = 0
)
```

Arguments

f1	Numeric or complex vector with factor 1 levels
f2	Numeric or complex vector with factor 2 levels
response	Numerical vector containing the response of the experiment.
norm	Error normality test (default is Shapiro-Wilk)
homog	Homogeneity test of variances (default is Bartlett)
alpha.f	Level of significance of the F test (default is 0.05)
alpha.t	Significance level of the multiple comparison test (<i>default</i> is 0.05)
quali	Defines whether the factor is quantitative or qualitative (qualitative)
names.fat	Name of factors
mcomp	Multiple comparison test (Tukey (default), LSD, Scott-Knott and Duncan)
grau	Polynomial degree in case of quantitative factor ($default$ is 1). Provide a vector with two elements.
grau12	Polynomial degree in case of quantitative factor ($default$ is 1). Provide a vector with n levels of factor 2, in the case of interaction f1 x f2 and qualitative factor 2 and quantitative factor 1.
grau21	Polynomial degree in case of quantitative factor ($default$ is 1). Provide a vector with n levels of factor 1, in the case of interaction f1 x f2 and qualitative factor 1 and quantitative factor 2.

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transf Applies data transformation (default is 1; for log consider 0; 'angular' for angu-

lar transformation)

constant Add a constant for transformation (enter value)

geom Graph type (columns or segments (For simple effect only))

theme ggplot2 theme (*default* is theme_classic())

ylab Variable response name (Accepts the *expression*() function) xlab Treatments name (Accepts the *expression*() function)

xlab. factor Provide a vector with two observations referring to the x-axis name of factors 1

and 2, respectively, when there is an isolated effect of the factors. This argument

uses 'parse'.

legend Legend title name

color Column chart color (default is "rainbow")

fill Defines chart color (to generate different colors for different treatments, define

fill = "trat")

textsize Font size labelsize Label Size

addmean Plot the average value on the graph (*default* is TRUE)

errorbar Plot the standard deviation bar on the graph (In the case of a segment and column

graph) - default is TRUE

CV Plotting the coefficient of variation and p-value of Anova (*default* is TRUE)

dec Number of cells

width.column Width column if geom="bar"

width.bar Width errorbar

angle x-axis scale text rotation

posi Legend position family Font family

point This function defines whether the point must have all points ("all"), mean ("mean"),

standard deviation (*default* - "mean_sd") or mean with standard error ("mean_se") if quali= FALSE. For quali=TRUE, 'mean_sd' and 'mean_se' change which in-

formation will be displayed in the error bar.

sup Number of units above the standard deviation or average bar on the graph

ylim y-axis scale angle.label Label angle

Value

The table of analysis of variance, the test of normality of errors (Shapiro-Wilk, Lilliefors, Anderson-Darling, Cramer-von Mises, Pearson and Shapiro-Francia), the test of homogeneity of variances (Bartlett or Levene), the test of independence of Durbin-Watson errors, the test of multiple comparisons (Tukey, LSD, Scott-Knott or Duncan) or adjustment of regression models up to grade 3 polynomial, in the case of quantitative treatments. The column chart for qualitative treatments is also returned.

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Note

The order of the chart follows the alphabetical pattern. Please use 'scale_x_discrete' from package ggplot2, 'limits' argument to reorder x-axis. The bars of the column and segment graphs are standard deviation.

The function does not perform multiple regression in the case of two quantitative factors.

In the final output when transformation (transf argument) is different from 1, the columns resp and respo in the mean test are returned, indicating transformed and non-transformed mean, respectively.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Leandro Simoes Azeredo Goncalves

Rodrigo Yudi Palhaci Marubayashi

References

Principles and procedures of statistics a biometrical approach Steel & Torry & Dickey. Third Edition 1997

Multiple comparisons theory and methods. Departament of statistics the Ohio State University. USA, 1996. Jason C. Hsu. Chapman Hall/CRC.

Practical Nonparametrics Statistics. W.J. Conover, 1999

Ramalho M.A.P., Ferreira D.F., Oliveira A.C. 2000. Experimentacao em Genetica e Melhoramento de Plantas. Editora UFLA.

Scott R.J., Knott M. 1974. A cluster analysis method for grouping mans in the analysis of variance. Biometrics, 30, 507-512.

Mendiburu, F., & de Mendiburu, M. F. (2019). Package 'agricolae'. R Package, Version, 1-2.

See Also

FAT2DIC.ad

Examples

FAT2DIC.ad 75

```
with(corn, FAT2DIC(A, B, Resp, mcomp="sk", quali=c(TRUE, TRUE),ylab="Heigth (cm)"))
```

FAT2DIC.ad

Analysis: DIC experiment in double factorial design with an additional treatment

Description

Analysis of an experiment conducted in a completely randomized design in a double factorial scheme using analysis of variance of fixed effects.

```
FAT2DIC.ad(
  f1,
  f2,
  repe,
  response,
  responseAd,
  norm = "sw",
  homog = "bt",
  alpha.f = 0.05,
  alpha.t = 0.05,
  quali = c(TRUE, TRUE),
  names.fat = c("F1", "F2"),
 mcomp = "tukey",
  grau = c(NA, NA),
  grau12 = NA,
  grau21 = NA,
  transf = 1,
  constant = 0,
  geom = "bar",
  theme = theme_classic(),
  ylab = "Response",
  xlab = "",
  xlab.factor = c("F1", "F2"),
  legend = "Legend",
  ad.label = "Additional",
  color = "rainbow",
  fill = "lightblue",
  textsize = 12,
  labelsize = 4,
  addmean = TRUE,
  errorbar = TRUE,
  CV = TRUE,
  dec = 3,
```

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```
width.column = 0.9,
width.bar = 0.3,
angle = 0,
posi = "right",
family = "sans",
point = "mean_sd",
sup = NA,
ylim = NA,
angle.label = 0
```

Arguments

f1 Numeric or complex vector with factor 1 levels
f2 Numeric or complex vector with factor 2 levels
repe Numeric or complex vector with repetitions

response Numerical vector containing the response of the experiment.

responseAd Numerical vector with additional treatment responses

norm Error normality test (default is Shapiro-Wilk)
homog Homogeneity test of variances (default is Bartlett)
alpha.f Level of significance of the F test (default is 0.05)

alpha.t Significance level of the multiple comparison test (*default* is 0.05) quali Defines whether the factor is quantitative or qualitative (*qualitative*)

names.fat Name of factors

mcomp Multiple comparison test (Tukey (*default*), LSD and Duncan)

grau Polynomial degree in case of quantitative factor (*default* is 1). Provide a vector

with two elements.

grau12 Polynomial degree in case of quantitative factor (*default* is 1). Provide a vector

with n levels of factor 2, in the case of interaction f1 x f2 and qualitative factor

2 and quantitative factor 1.

grau21 Polynomial degree in case of quantitative factor (*default* is 1). Provide a vector

with n levels of factor 1, in the case of interaction f1 x f2 and qualitative factor

1 and quantitative factor 2.

transf Applies data transformation (default is 1; for log consider 0; 'angular' for angu-

lar transformation)

constant Add a constant for transformation (enter value)

geom Graph type (columns or segments (For simple effect only))

theme ggplot2 theme (default is theme_classic())

ylab Variable response name (Accepts the *expression*() function)

xlab Treatments name (Accepts the *expression*() function)

xlab.factor Provide a vector with two observations referring to the x-axis name of factors 1

and 2, respectively, when there is an isolated effect of the factors. This argument

uses 'parse'.

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legend Legend title name ad.label Aditional label

color Column chart color (default is "rainbow")

fill Defines chart color (to generate different colors for different treatments, define

fill = "trat")

textsize Font size labelsize Label Size

addmean Plot the average value on the graph (*default* is TRUE)

errorbar Plot the standard deviation bar on the graph (In the case of a segment and column

graph) - default is TRUE

CV Plotting the coefficient of variation and p-value of Anova (*default* is TRUE)

dec Number of cells

width.column Width column if geom="bar"

width.bar Width errorbar

angle x-axis scale text rotation

posi legend position family Font family

point This function defines whether the point must have all points ("all"), mean ("mean"),

standard deviation (*default* - "mean_sd") or mean with standard error ("mean_se") if quali= FALSE. For quali=TRUE, 'mean_sd' and 'mean_se' change which in-

formation will be displayed in the error bar.

sup Number of units above the standard deviation or average bar on the graph

ylim y-axis scale angle.label label angle

Value

The table of analysis of variance, the test of normality of errors (Shapiro-Wilk, Lilliefors, Anderson-Darling, Cramer-von Mises, Pearson and Shapiro-Francia), the test of homogeneity of variances (Bartlett or Levene), the test of independence of Durbin-Watson errors, the test of multiple comparisons (Tukey, LSD, Scott-Knott or Duncan) or adjustment of regression models up to grade 3 polynomial, in the case of quantitative treatments. The column chart for qualitative treatments is also returned.

Note

The order of the chart follows the alphabetical pattern. Please use 'scale_x_discrete' from package ggplot2, 'limits' argument to reorder x-axis. The bars of the column and segment graphs are standard deviation.

The function does not perform multiple regression in the case of two quantitative factors.

The assumptions of variance analysis disregard additional treatment

In the final output when transformation (transf argument) is different from 1, the columns resp and respo in the mean test are returned, indicating transformed and non-transformed mean, respectively.

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Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi

References

Principles and procedures of statistics a biometrical approach Steel & Torry & Dickey. Third Edition 1997

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Practical Nonparametrics Statistics. W.J. Conover, 1999

Ramalho M.A.P., Ferreira D.F., Oliveira A.C. 2000. Experimentacao em Genetica e Melhoramento de Plantas. Editora UFLA.

Scott R.J., Knott M. 1974. A cluster analysis method for grouping mans in the analysis of variance. Biometrics, 30, 507-512.

Mendiburu, F., & de Mendiburu, M. F. (2019). Package 'agricolae'. R Package, Version, 1-2.

See Also

FAT2DIC

dunnett

Examples

```
library(AgroR)
data(cloro)
respAd=c(268, 322, 275, 350, 320)
with(cloro, FAT2DIC.ad(f1, f2, bloco, resp, respAd, ylab="Number of nodules", legend = "Stages"))
```

fat2_table

Utils: Summary of the analysis for factor arrangement with two qualitative factors.

Description

Summarizes the output returned in the summarise_anova function in list form. The advantage is that the table, in the case of significant interaction, is returned in a format that facilitates assembly in terms of scientific publication.

```
fat2_table(output, nf1, nf2, column = 1)
```

Arguments

output	Output of summarise_anova function for FAT2DIC, FAT2DIC.ad, FAT2DBC,
	FAT2DBC.ad, PSUBDIC and PSUBDBC design.
nf1	Number of levels of factor 1

nf1 Number of levels of factor 1 nf2 Number of levels of factor 2

column Variable column

Value

returns a list containing analysis output for experiments in FAT2DIC, FAT2DIC.ad, FAT2DBC, FAT2DBC.ad, PSUBDIC and PSUBDBC design.

Author(s)

Gabriel Danilo Shimizu

Examples

FAT3DBC

Analysis: DBC experiments in triple factorial

Description

Analysis of an experiment conducted in a randomized block design in a triple factorial scheme using analysis of variance of fixed effects.

```
FAT3DBC(
    f1,
    f2,
    f3,
```

```
block,
  response,
  norm = "sw",
  alpha.f = 0.05,
  alpha.t = 0.05,
  quali = c(TRUE, TRUE, TRUE),
 mcomp = "tukey",
  transf = 1,
  constant = 0,
  names.fat = c("F1", "F2", "F3"),
 ylab = "Response",
 xlab = "",
 xlab.factor = c("F1", "F2", "F3"),
  sup = NA,
  grau = c(NA, NA, NA),
  grau12 = NA,
  grau13 = NA,
  grau23 = NA,
 grau21 = NA,
 grau31 = NA,
 grau32 = NA,
 grau123 = NA,
 grau213 = NA,
 grau312 = NA,
 fill = "lightblue",
  theme = theme_classic(),
  angulo = 0,
  errorbar = TRUE,
  addmean = TRUE,
  family = "sans",
  dec = 3,
  geom = "bar",
  textsize = 12,
  labelsize = 4,
 point = "mean_sd",
 angle.label = 0
)
```

Arguments

f1	Numeric or complex vector with factor 1 levels
f2	Numeric or complex vector with factor 2 levels
f3	Numeric or complex vector with factor 3 levels
block	Numerical or complex vector with blocks
response	Numerical vector containing the response of the experiment.
norm	Error normality test (default is Shapiro-Wilk)
alpha.f	Level of significance of the F test (default is 0.05)

alpha.t	Significance level of the multiple comparison test (default is 0.05)
quali	Defines whether the factor is quantitative or qualitative (qualitative)
mcomp	Multiple comparison test (Tukey (default), LSD, Scott-Knott and Duncan)
transf	Applies data transformation (<i>default</i> is 1; for log consider 0; 'angular' for angular transformation)
constant	Add a constant for transformation (enter value)
names.fat	Allows labeling the factors 1, 2 and 3.
ylab	Variable response name (Accepts the expression() function)
xlab	Treatments name (Accepts the expression() function)
xlab.factor	Provide a vector with two observations referring to the x-axis name of factors 1, 2 and 3, respectively, when there is an isolated effect of the factors. This argument uses 'parse'.
sup	Number of units above the standard deviation or average bar on the graph
grau	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with three elements.
grau12	Polynomial degree in case of quantitative factor ($default$ is 1). Provide a vector with n levels of factor 2, in the case of interaction f1 x f2 and qualitative factor 2 and quantitative factor 1.
grau13	Polynomial degree in case of quantitative factor ($default$ is 1). Provide a vector with n levels of factor 3, in the case of interaction f1 x f3 and qualitative factor 3 and quantitative factor 1.
grau23	Polynomial degree in case of quantitative factor ($default$ is 1). Provide a vector with n levels of factor 3, in the case of interaction f2 x f3 and qualitative factor 3 and quantitative factor 2.
grau21	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 1, in the case of interaction f1 x f2 and qualitative factor 1 and quantitative factor 2.
grau31	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 1, in the case of interaction f1 x f3 and qualitative factor 1 and quantitative factor 3.
grau32	Polynomial degree in case of quantitative factor ($default$ is 1). Provide a vector with n levels of factor 2, in the case of interaction f2 x f3 and qualitative factor 2 and quantitative factor 3.
grau123	Polynomial degree in case of quantitative factor ($default$ is 1). Provide a vector with n levels of factor 1, in the case of interaction f1 x f2 x f3 and quantitative factor 1.
grau213	Polynomial degree in case of quantitative factor ($default$ is 1). Provide a vector with n levels of factor 2, in the case of interaction f1 x f2 x f3 and quantitative factor 2.
grau312	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 3, in the case of interaction $f1 \times f2 \times f3$ and quantitative factor 3.

fill Defines chart color (to generate different colors for different treatments, define

fill = "trat")

theme ggplot2 theme (default is theme_classic())

angulo x-axis scale text rotation

errorbar Plot the standard deviation bar on the graph (In the case of a segment and column

graph) - default is TRUE

addmean Plot the average value on the graph (*default* is TRUE)

family Font family dec Number of cells

geom Graph type (columns or segments)

textsize Font size labelsize Label Size

point This function defines whether the point must have all points ("all"), mean ("mean"),

standard deviation (*default* - "mean_sd") or mean with standard error ("mean_se") if quali= FALSE. For quali=TRUE, 'mean_sd' and 'mean_se' change which in-

formation will be displayed in the error bar.

angle.label label angle

Value

The analysis of variance table, the Shapiro-Wilk error normality test, the Bartlett homogeneity test of variances, the Durbin-Watson error independence test, multiple comparison test (Tukey, LSD, Scott-Knott or Duncan) or adjustment of regression models up to grade 3 polynomial, in the case of quantitative treatments. The column chart for qualitative treatments is also returned. For significant triple interaction only, no graph is returned.

Note

The order of the chart follows the alphabetical pattern. Please use 'scale_x_discrete' from package ggplot2, 'limits' argument to reorder x-axis. The bars of the column and segment graphs are standard deviation.

The function does not perform multiple regression in the case of two or more quantitative factors. The bars of the column and segment graphs are standard deviation.

In the final output when transformation (transf argument) is different from 1, the columns resp and respo in the mean test are returned, indicating transformed and non-transformed mean, respectively.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Leandro Simoes Azeredo Goncalves

Rodrigo Yudi Palhaci Marubayashi

References

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Practical Nonparametrics Statistics. W.J. Conover, 1999

Ramalho M.A.P., Ferreira D.F., Oliveira A.C. 2000. Experimentacao em Genetica e Melhoramento de Plantas. Editora UFLA.

Scott R.J., Knott M. 1974. A cluster analysis method for grouping mans in the analysis of variance. Biometrics, 30, 507-512.

Ferreira, E. B., Cavalcanti, P. P., and Nogueira, D. A. (2014). ExpDes: an R package for ANOVA and experimental designs. Applied Mathematics, 5(19), 2952.

Mendiburu, F., and de Mendiburu, M. F. (2019). Package 'agricolae'. R Package, Version, 1-2.

Examples

```
library(AgroR)
data(enxofre)
with(enxofre, FAT3DBC(f1, f2, f3, bloco, resp))
```

FAT3DBC.ad

Analysis: DBC experiments in triple factorial with aditional

Description

Analysis of an experiment conducted in a randomized block design in a triple factorial scheme with one aditional control using analysis of variance of fixed effects.

```
FAT3DBC.ad(
   f1,
   f2,
   f3,
   block,
   response,
   responseAd,
   norm = "sw",
   alpha.f = 0.05,
   alpha.t = 0.05,
   quali = c(TRUE, TRUE, TRUE),
   mcomp = "tukey",
   transf = 1,
   constant = 0,
   names.fat = c("F1", "F2", "F3"),
```

```
ylab = "Response",
 xlab = "",
 xlab.factor = c("F1", "F2", "F3"),
  sup = NA,
 grau = c(NA, NA, NA),
 grau12 = NA,
 grau13 = NA,
 grau23 = NA,
 grau21 = NA,
 grau31 = NA,
 grau32 = NA,
 grau123 = NA,
 grau213 = NA,
 grau312 = NA,
 fill = "lightblue",
  theme = theme_classic(),
  ad.label = "Additional",
  angulo = 0,
 errorbar = TRUE,
  addmean = TRUE,
  family = "sans",
 dec = 3,
 geom = "bar",
  textsize = 12,
 labelsize = 4,
 point = "mean_sd",
 angle.label = 0
)
```

Arguments

f1	Numeric or complex vector with factor 1 levels
f2	Numeric or complex vector with factor 2 levels
f3	Numeric or complex vector with factor 3 levels
block	Numerical or complex vector with blocks
response	Numerical vector containing the response of the experiment.
responseAd	Numerical vector containing the aditional response
norm	Error normality test (default is Shapiro-Wilk)
alpha.f	Level of significance of the F test (<i>default</i> is 0.05)
alpha.t	Significance level of the multiple comparison test (default is 0.05)
quali	Defines whether the factor is quantitative or qualitative (qualitative)
mcomp	Multiple comparison test (Tukey (default), LSD, Scott-Knott and Duncan)
transf	Applies data transformation (<i>default</i> is 1; for log consider 0; 'angular' for angular transformation)
constant	Add a constant for transformation (enter value)

Cat	Aller a lel eller de Control 1, 2 en 1,2
names.fat	Allows labeling the factors 1, 2 and 3.
ylab	Variable response name (Accepts the <i>expression</i> () function)
xlab	Treatments name (Accepts the <i>expression</i> () function)
xlab.factor	Provide a vector with two observations referring to the x-axis name of factors 1, 2 and 3, respectively, when there is an isolated effect of the factors. This argument uses 'parse'.
sup	Number of units above the standard deviation or average bar on the graph
grau	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with three elements.
grau12	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 2, in the case of interaction $f1 \times f2$ and qualitative factor 2 and quantitative factor 1.
grau13	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 3, in the case of interaction $f1 \times f3$ and qualitative factor 3 and quantitative factor 1.
grau23	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 3, in the case of interaction $f2 \times f3$ and qualitative factor 3 and quantitative factor 2.
grau21	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 1, in the case of interaction $f1 \times f2$ and qualitative factor 1 and quantitative factor 2.
grau31	Polynomial degree in case of quantitative factor ($default$ is 1). Provide a vector with n levels of factor 1, in the case of interaction f1 x f3 and qualitative factor 1 and quantitative factor 3.
grau32	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 2, in the case of interaction $f2 \times f3$ and qualitative factor 2 and quantitative factor 3.
grau123	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 1, in the case of interaction $f1 \times f2 \times f3$ and quantitative factor 1.
grau213	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 2, in the case of interaction $f1 \times f2 \times f3$ and quantitative factor 2.
grau312	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 3, in the case of interaction $f1 \times f2 \times f3$ and quantitative factor 3.
fill	Defines chart color (to generate different colors for different treatments, define fill = "trat")
theme	ggplot2 theme (default is theme_classic())
ad.label	Aditional label
angulo	x-axis scale text rotation
errorbar	Plot the standard deviation bar on the graph (In the case of a segment and column graph) - <i>default</i> is TRUE

addmean Plot the average value on the graph (*default* is TRUE)

family Font family dec Number of cells

geom Graph type (columns or segments)

textsize Font size labelsize Label size

point This function defines whether the point must have all points ("all"), mean ("mean"),

standard deviation (*default* - "mean_sd") or mean with standard error ("mean_se") if quali= FALSE. For quali=TRUE, 'mean_sd' and 'mean_se' change which in-

formation will be displayed in the error bar.

angle.label label angle

Value

The analysis of variance table, the Shapiro-Wilk error normality test, the Bartlett homogeneity test of variances, the Durbin-Watson error independence test, multiple comparison test (Tukey, LSD, Scott-Knott or Duncan) or adjustment of regression models up to grade 3 polynomial, in the case of quantitative treatments. The column chart for qualitative treatments is also returned. For significant triple interaction only, no graph is returned.

Note

The order of the chart follows the alphabetical pattern. Please use 'scale_x_discrete' from package ggplot2, 'limits' argument to reorder x-axis. The bars of the column and segment graphs are standard deviation.

The function does not perform multiple regression in the case of two or more quantitative factors. The bars of the column and segment graphs are standard deviation.

In the final output when transformation (transf argument) is different from 1, the columns resp and respo in the mean test are returned, indicating transformed and non-transformed mean, respectively.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Leandro Simoes Azeredo Goncalves

Rodrigo Yudi Palhaci Marubayashi

References

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Ferreira, E. B., Cavalcanti, P. P., and Nogueira, D. A. (2014). ExpDes: an R package for ANOVA and experimental designs. Applied Mathematics, 5(19), 2952.

Mendiburu, F., and de Mendiburu, M. F. (2019). Package 'agricolae'. R Package, Version, 1-2.

Examples

```
library(AgroR)
data(enxofre)
respAd=c(2000,2400,2530,2100)
attach(enxofre)
with(enxofre, FAT3DBC.ad(f1, f2, f3, bloco, resp, respAd))
```

FAT3DIC

Analysis: DIC experiments in triple factorial

Description

Analysis of an experiment conducted in a completely randomized design in a triple factorial scheme using analysis of variance of fixed effects.

```
FAT3DIC(
  f1,
  f2,
  f3,
  response,
  norm = "sw",
  alpha.t = 0.05,
  alpha.f = 0.05,
  quali = c(TRUE, TRUE, TRUE),
 mcomp = "tukey",
  grau = c(NA, NA, NA),
  grau12 = NA,
  grau13 = NA,
  grau23 = NA,
  grau21 = NA,
  grau31 = NA,
  grau32 = NA,
  grau123 = NA,
  grau213 = NA,
  grau312 = NA,
  transf = 1,
  constant = 0,
  names.fat = c("F1", "F2", "F3"),
```

```
ylab = "Response",
 xlab = "",
 xlab.factor = c("F1", "F2", "F3"),
  sup = NA,
 fill = "lightblue",
  theme = theme_classic(),
  angulo = 0,
 family = "sans",
  addmean = TRUE,
 errorbar = TRUE,
 dec = 3,
 geom = "bar",
  textsize = 12,
 labelsize = 4,
 point = "mean_sd",
 angle.label = 0
)
```

Arguments

f1	Numeric or complex vector with factor 1 levels
f2	Numeric or complex vector with factor 2 levels
f3	Numeric or complex vector with factor 3 levels
response	Numerical vector containing the response of the experiment.
norm	Error normality test (default is Shapiro-Wilk)
alpha.t	Significance level of the multiple comparison test (default is 0.05)
alpha.f	Level of significance of the F test (default is 0.05)
quali	Defines whether the factor is quantitative or qualitative (qualitative)
mcomp	Multiple comparison test (Tukey (default), LSD, Scott-Knott and Duncan)
grau	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with three elements.
grau12	Polynomial degree in case of quantitative factor ($default$ is 1). Provide a vector with n levels of factor 2, in the case of interaction f1 x f2 and qualitative factor 2 and quantitative factor 1.
grau13	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 3, in the case of interaction $f1 \times f3$ and qualitative factor 3 and quantitative factor 1.
grau23	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 3, in the case of interaction $f2 \times f3$ and qualitative factor 3 and quantitative factor 2.
grau21	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 1, in the case of interaction f1 x f2 and qualitative factor 1 and quantitative factor 2.

grau31	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 1, in the case of interaction f1 x f3 and qualitative factor 1 and quantitative factor 3.
grau32	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 2, in the case of interaction f2 x f3 and qualitative factor 2 and quantitative factor 3.
grau123	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 1, in the case of interaction f1 x f2 x f3 and quantitative factor 1.
grau213	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 2, in the case of interaction f1 x f2 x f3 and quantitative factor 2.
grau312	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 3, in the case of interaction f1 x f2 x f3 and quantitative factor 3.
transf	Applies data transformation (<i>default</i> is 1; for log consider 0; 'angular' for angular transformation)
constant	Add a constant for transformation (enter value)
names.fat	Allows labeling the factors 1, 2 and 3.
ylab	Variable response name (Accepts the expression() function)
xlab	treatments name (Accepts the expression() function)
xlab.factor	Provide a vector with two observations referring to the x-axis name of factors 1, 2 and 3, respectively, when there is an isolated effect of the factors. This argument uses 'parse'.
sup	Number of units above the standard deviation or average bar on the graph
fill	Defines chart color (to generate different colors for different treatments, define fill = "trat")
theme	ggplot2 theme (default is theme_classic())
angulo	x-axis scale text rotation
family	Font family
addmean	Plot the average value on the graph (default is TRUE)
errorbar	Plot the standard deviation bar on the graph (In the case of a segment and column graph) - <i>default</i> is TRUE
dec	Number of cells
geom	Graph type (columns or segments)
textsize	Font size
labelsize	Label Size
point	This function defines whether the point must have all points ("all"), mean ("mean"), standard deviation (<i>default</i> - "mean_sd") or mean with standard error ("mean_se") if quali= FALSE. For quali=TRUE, 'mean_sd' and 'mean_se' change which information will be displayed in the error bar.
angle.label	label angle

Value

The analysis of variance table, the Shapiro-Wilk error normality test, the Bartlett homogeneity test of variances, the Durbin-Watson error independence test, multiple comparison test (Tukey, LSD, Scott-Knott or Duncan) or adjustment of regression models up to grade 3 polynomial, in the case of quantitative treatments. The column chart for qualitative treatments is also returned. For significant triple interaction only, no graph is returned.

Note

The order of the chart follows the alphabetical pattern. Please use 'scale_x_discrete' from package ggplot2, 'limits' argument to reorder x-axis. The bars of the column and segment graphs are standard deviation.

The function does not perform multiple regression in the case of two or more quantitative factors. The bars of the column and segment graphs are standard deviation.

In the final output when transformation (transf argument) is different from 1, the columns resp and respo in the mean test are returned, indicating transformed and non-transformed mean, respectively.

Author(s)

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Mendiburu, F., and de Mendiburu, M. F. (2019). Package 'agricolae'. R Package, Version, 1-2.

Examples

```
library(AgroR)
data(enxofre)
with(enxofre, FAT3DIC(f1, f2, f3, resp))
```

FAT3DIC.ad

Analysis: DIC experiments in triple factorial with aditional

Description

Analysis of an experiment conducted in a completely randomized design in a triple factorial scheme with one aditional control using analysis of variance of fixed effects.

```
FAT3DIC.ad(
  f1,
  f2,
  f3,
  repe,
  response,
  responseAd,
  norm = "sw",
  alpha.f = 0.05,
  alpha.t = 0.05,
  quali = c(TRUE, TRUE, TRUE),
  mcomp = "tukey",
  transf = 1,
  constant = 0,
  names.fat = c("F1", "F2", "F3"),
  ylab = "Response",
  xlab = "",
  xlab.factor = c("F1", "F2", "F3"),
  sup = NA,
  grau = c(NA, NA, NA),
  grau12 = NA,
  grau13 = NA,
  grau23 = NA,
  grau21 = NA,
  grau31 = NA,
  grau32 = NA,
  grau123 = NA,
  grau213 = NA,
  grau312 = NA,
  fill = "lightblue",
  theme = theme_classic(),
  ad.label = "Additional",
  angulo = 0,
  errorbar = TRUE,
  addmean = TRUE,
  family = "sans",
  dec = 3,
```

```
geom = "bar",
textsize = 12,
labelsize = 4,
point = "mean_sd",
angle.label = 0
)
```

Arguments

Numeric or complex vector with factor 1 levels
Numeric or complex vector with factor 2 levels
Numeric or complex vector with factor 3 levels

repe Numerical or complex vector with blocks

response Numerical vector containing the response of the experiment.

responseAd Numerical vector containing the aditional response norm Error normality test (*default* is Shapiro-Wilk)

alpha.f Level of significance of the F test (*default* is 0.05)

alpha.t Significance level of the multiple comparison test (*default* is 0.05) quali Defines whether the factor is quantitative or qualitative (*qualitative*)

mcomp Multiple comparison test (Tukey (default), LSD, Scott-Knott and Duncan)

transf Applies data transformation (default is 1; for log consider 0; 'angular' for angu-

lar transformation)

constant Add a constant for transformation (enter value)

names.fat Allows labeling the factors 1, 2 and 3.

ylab Variable response name (Accepts the *expression*() function)

xlab Treatments name (Accepts the *expression*() function)

xlab. factor Provide a vector with two observations referring to the x-axis name of factors

1, 2 and 3, respectively, when there is an isolated effect of the factors. This

argument uses 'parse'.

sup Number of units above the standard deviation or average bar on the graph

grau Polynomial degree in case of quantitative factor (default is 1). Provide a vector

with three elements.

grau12 Polynomial degree in case of quantitative factor (*default* is 1). Provide a vector

with n levels of factor 2, in the case of interaction f1 x f2 and qualitative factor

2 and quantitative factor 1.

grau13 Polynomial degree in case of quantitative factor (*default* is 1). Provide a vector

with n levels of factor 3, in the case of interaction f1 x f3 and qualitative factor

3 and quantitative factor 1.

grau23 Polynomial degree in case of quantitative factor (*default* is 1). Provide a vector

with n levels of factor 3, in the case of interaction f2 x f3 and qualitative factor

3 and quantitative factor 2.

grau21	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 1, in the case of interaction f1 x f2 and qualitative factor 1 and quantitative factor 2.
grau31	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 1, in the case of interaction f1 x f3 and qualitative factor 1 and quantitative factor 3.
grau32	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 2, in the case of interaction f2 x f3 and qualitative factor 2 and quantitative factor 3.
grau123	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 1, in the case of interaction f1 x f2 x f3 and quantitative factor 1.
grau213	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 2, in the case of interaction f1 x f2 x f3 and quantitative factor 2.
grau312	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 3, in the case of interaction f1 x f2 x f3 and quantitative factor 3.
fill	Defines chart color (to generate different colors for different treatments, define fill = "trat")
theme	ggplot2 theme (default is theme_classic())
ad.label	Aditional label
angulo	x-axis scale text rotation
errorbar	Plot the standard deviation bar on the graph (In the case of a segment and column graph) - <i>default</i> is TRUE
addmean	Plot the average value on the graph (default is TRUE)
family	Font family
dec	Number of cells
geom	Graph type (columns or segments)
textsize	Font size
labelsize	Label size
point	This function defines whether the point must have all points ("all"), mean ("mean"), standard deviation (<i>default</i> - "mean_sd") or mean with standard error ("mean_se") if quali= FALSE. For quali=TRUE, 'mean_sd' and 'mean_se' change which information will be displayed in the error bar.
angle.label	label angle

Value

The analysis of variance table, the Shapiro-Wilk error normality test, the Bartlett homogeneity test of variances, the Durbin-Watson error independence test, multiple comparison test (Tukey, LSD, Scott-Knott or Duncan) or adjustment of regression models up to grade 3 polynomial, in the case of quantitative treatments. The column chart for qualitative treatments is also returned. For significant triple interaction only, no graph is returned.

Note

The order of the chart follows the alphabetical pattern. Please use 'scale_x_discrete' from package ggplot2, 'limits' argument to reorder x-axis. The bars of the column and segment graphs are standard deviation.

The function does not perform multiple regression in the case of two or more quantitative factors. The bars of the column and segment graphs are standard deviation.

In the final output when transformation (transf argument) is different from 1, the columns resp and respo in the mean test are returned, indicating transformed and non-transformed mean, respectively.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Leandro Simoes Azeredo Goncalves

Rodrigo Yudi Palhaci Marubayashi

References

Principles and procedures of statistics a biometrical approach Steel, Torry and Dickey. Third Edition 1997

Multiple comparisons theory and methods. Departament of statistics the Ohio State University. USA, 1996. Jason C. Hsu. Chapman Hall/CRC.

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Ramalho M.A.P., Ferreira D.F., Oliveira A.C. 2000. Experimentacao em Genetica e Melhoramento de Plantas. Editora UFLA.

Scott R.J., Knott M. 1974. A cluster analysis method for grouping mans in the analysis of variance. Biometrics, 30, 507-512.

Ferreira, E. B., Cavalcanti, P. P., and Nogueira, D. A. (2014). ExpDes: an R package for ANOVA and experimental designs. Applied Mathematics, 5(19), 2952.

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Examples

```
library(AgroR)
data(enxofre)
respAd=c(2000,2400,2530,2100)
with(enxofre, FAT3DIC.ad(f1, f2, f3, bloco, resp, respAd))
```

grid.onefactor 95

grid.onefactor	utils: group graphs of the output of simple experiments in dic, dbc or
	dql

Description

group graphs of the output of simple experiments into dic, dbc or dql. It is possible to group up to 6 graphs in different arrangements (see model argument)

Usage

```
grid.onefactor(output, model = "type1")
```

Arguments

output Vector with the outputs of the DIC, DBC or DQL functions

model Graph arrangement model, see in detail.

Details

- 'type1': Two graphs next to each other - 'type2': Two graphs one below the other - 'type3': Three graphs, two top and one centered below - 'type4': Three graphs one below the other - 'type5': Four graphs, two at the top and two at the bottom - 'type6': Four graphs one below the other - 'type7': Five graphs, two at the top, two in the middle and one centered at the bottom - 'type8': Five graphs, three at the top, two centered at the bottom - 'type9': Six graphs, three at the top, three centered at the bottom - 'type10': Six graphs, two at the top, two in the middle and two at the bottom

Value

returns grouped graphs

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

Examples

```
data("pomegranate")
attach(pomegranate)
a=DIC(trat, WL, geom = "point", ylab = "WL")
b=DIC(trat, SS, geom = "point", ylab="SS")
c=DIC(trat, AT, geom = "point", ylab = "AT")
grid.onefactor(c(a,b),model = "type1")
grid.onefactor(c(a,b),model = "type2")
grid.onefactor(c(a,b,c),model = "type3")
grid.onefactor(c(a,b,c),model = "type4")
```

96 jointcluster

ibarplot.double

Graph: Invert letters for two factor chart

Description

invert uppercase and lowercase letters in graph for factorial scheme the subdivided plot with significant interaction

Usage

```
ibarplot.double(analysis)
```

Arguments

analysis

FAT2DIC, FAT2DBC, PSUBDIC or PSUBDBC object

Value

Return column chart for two factors

Examples

```
data(covercrops)
attach(covercrops)
a=FAT2DBC(A, B, Bloco, Resp, ylab=expression("Yield"~(Kg~"100 m"^2)),
legend = "Cover crops",alpha.f = 0.3,family = "serif")
ibarplot.double(a)
```

jointcluster

Analysis: Method to evaluate similarity of experiments based on QM-

Description

This function presents a method to evaluate similarity of experiments based on a matrix of QMres of all against all. This is used as a measure of similarity and applied in clustering.

Usage

```
jointcluster(qmres, information = "matrix", method.cluster = "ward.D")
```

Arguments

qmres Vector containing mean squares of residuals or output from list DIC or DBC

function

information Option to choose the return type. 'matrix', 'bar' or 'cluster'

method.cluster Grouping method

laranja 97

Value

Returns a residual mean square ratio matrix, bar graph with ratios sorted in ascending order, or cluster analysis.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Examples

```
qmres=c(0.344429, 0.300542, 0.124833, 0.04531, 0.039571, 0.011812, 0.00519)
jointcluster(qmres,information = "cluster")
jointcluster(qmres,information = "matrix")
jointcluster(qmres,information = "bar")

data(mirtilo)
m=lapply(unique(mirtilo$exp),function(x){
    m=with(mirtilo[mirtilo$exp==x,],DBC(trat,bloco,resp))})
jointcluster(m)
```

laranja

Dataset: Orange plants under different rootstocks

Description

An experiment was conducted with the objective of studying the behavior of nine rootstocks for the Valencia orange tree. The data set refers to the 1973 evaluation (12 years old). The rootstocks are: T1: Tangerine Sunki; T2: National rough lemon; T3: Florida rough lemon; T4: Cleopatra tangerine; T5: Citranger-troyer; T6: Trifoliata; T7: Clove Tangerine; T8: Country orange; T9: Clove Lemon. The number of fruits per plant was evaluated.

Usage

```
data(laranja)
```

Format

data.frame containing data set

f1 Categorical vector with treatments

bloco Categorical vector with block

resp Numeric vector with number of fruits per plant

References

Planejamento e Analise Estatistica de Experimentos Agronomicos (2013) - Decio Barbin - pg. 72

98 line_plot

See Also

cloro, enxofre, mirtilo, pomegranate, porco, sensorial, simulate1, simulate2, simulate3, tomate, weather, phao, passiflora, aristolochia

Examples

```
data(laranja)
```

line_plot

Graph: Line chart

Description

Performs a descriptive line graph with standard deviation bars

Usage

```
line_plot(
   time,
   response,
   factor = NA,
   errorbar = "sd",
   ylab = "Response",
   xlab = "Time",
   legend.position = "right",
   theme = theme_classic()
)
```

Arguments

time Vector containing the x-axis values
response Vector containing the y-axis values
factor Vector containing a categorical factor
errorbar Error bars (sd or se)

ylab y axis title

xlab x axis title

legend.position

Legend position

theme ggplot2 theme (default is theme_classic())

Value

Returns a line chart with error bars

logistic 99

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

See Also

```
radargraph, sk_graph, plot_TH, corgraph, spider_graph
```

Examples

```
dose=rep(c(0,2,4,6,8,10),e=3,2)
resp=c(seq(1,18,1),seq(2,19,1))
fator=rep(c("A","B"),e=18)
line_plot(dose,resp,fator)
```

logistic

Analysis: Logistic regression

Description

Logistic regression is a very popular analysis in agrarian sciences, such as in fruit growth curves, seed germination, etc...The logistic function performs the analysis using 3 or 4 parameters of the logistic model, being imported from the LL function .3 or LL.4 of the drc package (Ritz & Ritz, 2016).

```
logistic(
   trat,
   resp,
   npar = "LL.3",
   error = "SE",
   ylab = "Dependent",
   xlab = expression("Independent"),
   theme = theme_classic(),
   legend.position = "top",
   r2 = "all",
   width.bar = NA,
   scale = "none",
   textsize = 12,
   font.family = "sans"
)
```

100 logistic

Arguments

trat Numerical or complex vector with treatments

resp Numerical vector containing the response of the experiment.

npar Number of model parameters

error Error bar (It can be SE - default, SD or FALSE)

ylab Variable response name (Accepts the *expression*() function)

xlab Treatments name (Accepts the *expression*() function)

theme ggplot2 theme (default is theme_bw())

legend.position

Legend position (*default* is c(0.3,0.8))

r2 Coefficient of determination of the mean or all values (*default* is all)

width.bar Bar width

scale Sets x scale (*default* is none, can be "log")

textsize Font size

font.family Font family (*default* is sans)

Details

The three-parameter log-logistic function with lower limit 0 is

$$f(x) = 0 + \frac{d}{1 + \exp(b(\log(x) - \log(e)))}$$

The four-parameter log-logistic function is given by the expression

$$f(x) = c + \frac{d - c}{1 + \exp(b(\log(x) - \log(e)))}$$

The function is symmetric about the inflection point (e).

Value

The function allows the automatic graph and equation construction of the logistic model, provides important statistics, such as the Akaike (AIC) and Bayesian (BIC) inference criteria, coefficient of determination (r2), square root of the mean error (RMSE).

Author(s)

Model imported from the drc package (Ritz et al., 2016)

Gabriel Danilo Shimizu

Leandro Simoes Azeredo Goncalves

References

2016.

Seber, G. A. F. and Wild, C. J (1989) Nonlinear Regression, New York: Wiley and Sons (p. 330). Ritz, C.; Strebig, J.C.; Ritz, M.C. Package 'drc'. Creative Commons: Mountain View, CA, USA,

mirtilo 101

Examples

```
data("emerg")
with(emerg, logistic(time, resp,xlab="Time (days)",ylab="Emergence (%)"))
with(emerg, logistic(time, resp,npar="LL.4",xlab="Time (days)",ylab="Emergence (%)"))
```

mirtilo

Dataset: Cutting blueberry data

Description

An experiment was carried out in order to evaluate the rooting (resp1) of blueberry cuttings as a function of the cutting size (Treatment Colume). This experiment was repeated three times (Location column) and a randomized block design with four replications was adopted.

Usage

```
data(mirtilo)
```

Format

```
data.frame containing data set

trat Categorical vector with treatments
exp Categorical vector with experiment
bloco Categorical vector with block
resp Numeric vector
```

See Also

cloro, enxofre, laranja, pomegranate, porco, sensorial, simulate1, simulate2, simulate3, tomate, weather

Examples

```
data(mirtilo)
attach(mirtilo)
```

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orchard

Dataset: Orchard

Description

An experiment was carried out to analyze the treatments in orchards applied in the rows and between the rows, in a split-plot scheme according to a randomized block design. For this case, the line and leading are considered the levels of the factor applied in the plots and the treatments are considered the levels of the factor applied in the subplots. Microbial biomass carbon was analyzed.

Usage

data(orchard)

Format

data.frame containing data set

A Categorical vector with plot

B Categorical vector with split-plot

Bloco Categorical vector with block

Resp Numeric vector with microbial biomass carbon

See Also

enxofre, laranja, mirtilo, pomegranate, porco, sensorial, simulate1, simulate2, simulate3, tomate, weather, phao, passiflora, aristolochia

Examples

data(orchard)

passiflora

Dataset: Substrate data in the production of passion fruit seedlings

Description

An experiment was carried out in order to evaluate the influence of the substrate on the dry mass of aerial part and root in yellow sour passion fruit. The experiment was conducted in a randomized block design with four replications. The treatments consisted of five substrates (Vermiculite, MC Normal, Carolina Soil, Mc organic and sand)

Usage

data(passiflora)

PCA_function 103

Format

```
data.frame containing data set

trat Categorical vector with substrate

bloco Categorical vector with block

MSPA Numeric vector with dry mass of aerial part

MSR Numeric vector with dry mass of root
```

See Also

cloro, enxofre, laranja, mirtilo, pomegranate, porco, sensorial, simulate1, simulate2, simulate3, tomate, weather

Examples

```
data(passiflora)
```

PCA_function

Analysis: Principal components analysis

Description

This function performs principal component analysis.

```
PCA_function(
  data,
  scale = TRUE,
  text = TRUE,
  pointsize = 5,
  textsize = 12,
  labelsize = 4,
  linesize = 0.6,
  repel = TRUE,
  ylab = NA,
  xlab = NA,
  groups = NA,
  sc = 1,
  font.family = "sans",
  theme = theme_bw(),
  label.legend = "Cluster",
  type.graph = "biplot"
)
```

104 PCA_function

Arguments

linesize

data Data.frame with data set. Line name must indicate the treatment

scale Performs data standardization (*default* is TRUE)

text Add label (default is TRUE)
pointsize Point size (default is 5)
textsize Text size (default is 12)
labelsize Label size (default is 4)

repel Avoid text overlay (default is TRUE)

Line size (default is 0.8)

ylab Names y-axis
xlab Names x-axis
groups Define grouping

sc Secondary axis scale ratio (default is 1)

font.family Font family (*default* is sans)

theme Theme ggplot2 (default is theme_bw())
label.legend Legend title (when group is not NA)
type.graph Type of chart (default is biplot)

Details

The type.graph argument defines the graph that will be returned, in the case of "biplot" the biplot graph is returned with the first two main components and with eigenvalues and eigenvectors. In the case of "scores" only the treatment scores are returned, while for "cor" the correlations are returned. For "corPCA" a correlation between the vectors with the components is returned.

Value

The eigenvalues and eigenvectors, the explanation percentages of each principal component, the correlations between the vectors with the principal components, as well as graphs are returned.

Author(s)

Gabriel Danilo Shimizu

Examples

data(pomegranate)
medias=tabledesc(pomegranate)
PCA_function(medias)

pepper 105

pepper

Dataset: Pepper

Description

A vegetable breeder is characterizing five mini pepper accessions from the State University of Londrina germplasm bank for agronomic and biochemical variables. The experiment was conducted in a completely randomized design with four replications

Usage

```
data(pepper)
```

Format

data.frame containing data set

Acesso Categorical vector with accessions

MS Numeric vector com dry mass

VitC Numeric vector with Vitamin C

See Also

enxofre, laranja, mirtilo, pomegranate, porco, sensorial, simulate1, simulate2, simulate3, tomate, weather, phao, passiflora, aristolochia

Examples

data(pepper)

phao

Dataset: Osmocote in Phalaenopsis sp.

Description

The objective of the work was to evaluate the effect of doses of osmocote (15-09-12-N-P2O5-K2O, respectively) on the initial development of the orchid *Phalaenopsis* sp. The osmocote fertilizer was added in the following doses: 0, 2, 4, 6 and 8 g vase-1. After twelve months, leaf length was evaluated.

Usage

data(phao)

106 plot_cor

Format

```
data.frame containing data set

dose Numeric vector with doses

comp Numeric vector with leaf length
```

References

de Paula, J. C. B., Junior, W. A. R., Shimizu, G. D., Men, G. B., & de Faria, R. T. (2020). Fertilizante de liberacao controlada no crescimento inicial da orquidea *Phalaenopsis* sp. Revista Cultura Agronomica, 29(2), 289-299.

See Also

pomegranate, passiflora, cloro, enxofre, laranja, mirtilo, porco, sensorial, simulate1, simulate2, simulate3, tomate, weather

Examples

```
data(phao)
```

plot_cor

Graph: Plot correlation

Description

Correlation analysis function (Pearson or Spearman)

```
plot_cor(
 х,
 у,
 method = "pearson",
 ylab = "Dependent",
 xlab = "Independent",
  theme = theme_classic(),
  pointsize = 5,
  shape = 21,
  fill = "gray"
  color = "black",
  axis.size = 12,
  ic = TRUE,
  title = NA,
  family = "sans"
)
```

plot_interaction 107

Arguments

x Numeric vector with independent variable
 y Numeric vector with dependent variable
 method Method correlation (default is Pearson)

ylab Variable response name (Accepts the *expression*() function)

xlab Treatments name (Accepts the *expression*() function)

theme ggplot2 theme (default is theme_classic())

pointsize Point size
shape shape format
fill Fill point
color Color point
axis.size Axis text size

ic add interval of confidence

title title

family Font family

Value

The function returns a graph for correlation

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br> Leandro Simoes Azeredo Goncalves Rodrigo Yudi Palhaci Marubayashi

Examples

```
data("pomegranate")
with(pomegranate, plot_cor(WL, SS, xlab="WL", ylab="SS"))
```

plot_interaction Graph: Interaction plot

Description

Performs an interaction graph from an output of the FAT2DIC, FAT2DBC, PSUBDIC or PSUBDBC commands.

plot_interaction

Usage

```
plot_interaction(
   a,
   box_label = TRUE,
   repel = FALSE,
   pointsize = 3,
   linesize = 0.8,
   width.bar = 0.05,
   add.errorbar = TRUE
)
```

Arguments

a FAT2DIC, FAT2DBC, PSUBDIC or PSUBDBC object

box_label Add box in label

repel a boolean, whether to use ggrepel to avoid overplotting text labels or not.

pointsize Point size

linesize Line size (Trendline and Error Bar)

width.bar width of the error bars.

add.errorbar Add error bars.

Value

Returns an interaction graph with averages and letters from the multiple comparison test

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
```

Rodrigo Yudi Palhaci Marubayashi

Examples

```
data(cloro)
a=with(cloro, FAT2DIC(f1, f2, resp))
plot_interaction(a)
```

plot_jitter 109

plot_jitter

Graph: Column, box or segment chart with observations

Description

The function performs the construction of graphs of boxes, columns or segments with all the observations represented in the graph.

Usage

```
plot_jitter(model)
```

Arguments

model

DIC, DBC or DQL object

Value

Returns with graph of boxes, columns or segments with all the observations represented in the graph.

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

Examples

```
data("pomegranate")
a=with(pomegranate,DIC(trat,WL,geom="point"))
plot_jitter(a)
```

plot_TH

Graph: Climate chart of temperature and humidity

Description

The plot_TH function allows the user to build a column/line graph with climatic parameters of temperature (maximum, minimum and average) and relative humidity (UR) or precipitation. This chart is widely used in scientific work in agrarian science

plot_TH

Usage

```
plot_TH(
  tempo,
  Tmed,
  Tmax,
  Tmin,
 UR,
  xlab = "Time",
  yname1 = expression("Humidity (%)"),
  yname2 = expression("Temperature ("^o * "C)"),
  legend.H = "Humidity",
  legend.tmed = "Tmed",
  legend.tmin = "Tmin",
  legend.tmax = "Tmax",
  colormax = "red",
  colormin = "blue",
  colormean = "darkgreen",
  fillbar = "gray80",
  limitsy1 = c(0, 100),
  x = "days",
  breaks = "1 months",
  textsize = 12,
  legendsize = 12,
  titlesize = 12,
  linesize = 1,
  date_format = "%m-%Y",
  sc = 2.5,
  angle = 0,
  legend.position = "bottom",
  theme = theme_classic()
)
```

Arguments

tempo	Vector with times
Tmed	Vector with mean temperature
Tmax	Vector with maximum temperature
Tmin	Vector with minimum temperature
UR	Vector with relative humidity or precipitation
xlab	x axis name
yname1	y axis name
yname2	Secondary y-axis name
legend.H	Legend column
legend.tmed	Legend mean temperature
legend.tmin	Legend minimum temperature

plot_TH 111

legend. tmax Legend maximum temperature

colormax Maximum line color (default is "red")
colormin Minimum line color (default is "blue")
colormean Midline color (default is "darkgreen")
fillbar Column fill color (default is "gray80")
limitsy1 Primary y-axis scale (default is c(0,100))

x x scale type (days or data, default is "days")

breaks Range for x scale when x = "date" (default is 1 months)

textsize Axis text size
legendsize Legend text size
titlesize Axis title size
linesize Line size

date_format Date format for x="data"

sc Scale for secondary y-axis in relation to primary y-axis (declare the number of

times that y2 is less than or greater than y1, the default being 2.5)

angle x-axis scale text rotation

legend.position

Legend position

theme ggplot2 theme

Value

Returns row and column graphs for graphical representation of air temperature and relative humidity. Graph normally used in scientific articles

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br> Leandro Simoes Azeredo Goncalves Rodrigo Yudi Palhaci Marubayashi

See Also

radargraph, sk_graph, barplot_positive, corgraph, plot_TH1, spider_graph, line_plot

```
library(AgroR)
data(weather)
with(weather, plot_TH(tempo, Tmed, Tmax, Tmin, UR))
```

plot_TH1

plot_TH1

Graph: Climate chart of temperature and humidity (Model 2)

Description

The plot_TH1 function allows the user to build a column/line graph with climatic parameters of temperature (maximum, minimum and average) and relative humidity (UR) or precipitation. This chart is widely used in scientific work in agrarian science

Usage

```
plot_TH1(
  tempo,
  Tmed,
  Tmax,
  Tmin,
 UR,
  xlab = "Time",
 yname1 = expression("Humidity (%)"),
 yname2 = expression("Temperature ("^o * "C)"),
  legend.T = "Temperature",
  legend.H = "Humidity",
  legend.tmed = "Tmed",
  legend.tmin = "Tmin",
  legend.tmax = "Tmax",
  colormax = "red",
  colormin = "blue",
  colormean = "darkgreen",
  fillarea = "darkblue",
  facet.fill = "#FF9933",
  panel.grid = FALSE,
  x = "days",
  breaks = "1 months",
  textsize = 12,
  legendsize = 12,
  titlesize = 12,
  linesize = 1,
  date_format = "%m-%Y",
  angle = 0,
  legend.position = c(0.1, 0.3)
)
```

Arguments

tempo Vector with times

Tmed Vector with mean temperature

plot_TH1 113

Tmax Vector with maximum temperature
Tmin Vector with minimum temperature

UR Vector with relative humidity or precipitation

xlab x axis name yname1 y axis name

yname2 Secondary y-axis name legend.T faceted title legend 1 legend.H faceted title legend 2

legend.tmedLegend mean temperaturelegend.tminLegend minimum temperaturelegend.tmaxLegend maximum temperature

colormax Maximum line color (default is "red")
colormin Minimum line color (default is "blue")
colormean Midline color (default is "darkgreen")
fillarea area fill color (default is "darkblue")

facet.fill faceted title fill color (*default* is #FF9933)
panel.grid remove grid line (*default* is FALSE)

x x scale type (days or data, default is "days")

breaks Range for x scale when x = "date" (default is 1 months)

textsize Axis text size
legendsize Legend text size
titlesize Axis title size
linesize Line size

date_format Date format for x="data" angle x-axis scale text rotation

legend.position

Legend position

Value

Returns row and column graphs for graphical representation of air temperature and relative humidity. Graph normally used in scientific articles

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br> Leandro Simoes Azeredo Goncalves Rodrigo Yudi Palhaci Marubayashi

See Also

radargraph, sk_graph, barplot_positive, corgraph, spider_graph, line_plot

plot_tonetest

Examples

```
library(AgroR)
data(weather)
with(weather, plot_TH1(tempo, Tmed, Tmax, Tmin, UR))
```

plot_tonetest

Graphics: Graphic for t test to compare means with a reference value

Description

Sometimes the researcher wants to test whether the treatment mean is greater than/equal to or less than a reference value. For example, I want to know if the average productivity of my treatment is higher than the average productivity of a given country. For this, this function allows comparing the means with a reference value using the t test.

Usage

```
plot_tonetest(tonetest, alpha = 0.95)
```

Arguments

tonetest t.one.test object alpha confidence level.

Value

returns a density plot and a column plot to compare a reference value with other treatments.

Author(s)

Gabriel Danilo Shimizu

```
library(AgroR)
data("pomegranate")
resu=tonetest(resp=pomegranate$WL, trat=pomegranate$trat, mu=2)
plot_tonetest(resu)
```

polynomial 115

polynomial

Analysis: Linear regression graph

Description

Linear regression analysis of an experiment with a quantitative factor or isolated effect of a quantitative factor

Usage

```
polynomial(
  trat,
  resp,
 ylab = "Response",
 xlab = "Independent",
 yname.poly = "y",
  xname.poly = "x",
  grau = NA,
  theme = theme_classic(),
  point = "mean_sd",
  color = "gray80",
  posi = "top",
  textsize = 12,
  se = FALSE,
 ylim = NA,
  family = "sans",
  pointsize = 4.5,
  linesize = 0.8,
 width.bar = NA,
 n = NA,
 SSq = NA,
 DFres = NA
)
```

Arguments

trat	Numerical vector with treatments (Declare as numeric)
resp	Numerical vector containing the response of the experiment.
ylab	Dependent variable name (Accepts the expression() function)
xlab	Independent variable name (Accepts the <i>expression</i> () function)
yname.poly	Y name in equation
xname.poly	X name in equation
grau	Degree of the polynomial (1, 2 or 3)
theme	ggplot2 theme (<i>default</i> is theme_classic())

polynomial polynomial

point Defines whether to plot mean ("mean"), all repetitions ("all"), mean with stan-

dard deviation ("mean_sd") or mean with standard error (default - "mean_se").

color Graph color (default is gray80)

posi Legend position

textsize Font size

se Adds confidence interval (*default* is FALSE)

ylim y-axis scale
family Font family
pointsize Point size

linesize line size (Trendline and Error Bar)

width.bar width of the error bars of a regression graph.

n Number of decimal places for regression equations

SSq Sum of squares of the residue

DFres Residue freedom degrees

Value

Returns linear, quadratic or cubic regression analysis.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Leandro Simoes Azeredo Goncalves

Rodrigo Yudi Palhaci Marubayashi

See Also

```
polynomial2, polynomial2_color
```

```
data("phao")
with(phao, polynomial(dose,comp, grau = 2))
```

polynomial2 117

polynomial2

Analysis: Linear regression graph in double factorial

Description

Linear regression analysis for significant interaction of an experiment with two factors, one quantitative and one qualitative

Usage

```
polynomial2(
  fator1,
  resp,
 fator2,
 color = NA,
 grau = NA,
 ylab = "Response",
 xlab = "Independent",
  theme = theme_classic(),
  se = FALSE,
 point = "mean_sd",
 legend.title = "Treatments",
 posi = "top",
  textsize = 12,
 ylim = NA,
  family = "sans",
 width.bar = NA,
 pointsize = 3,
 linesize = 0.8,
 separate = c("(\"", "\")"),
 n = NA,
 DFres = NA,
  SSq = NA
)
```

Arguments

fator1	Numeric or complex vector with factor 1 levels
resp	Numerical vector containing the response of the experiment.
fator2	Numeric or complex vector with factor 2 levels
color	Graph color (default is NA)
grau	Degree of the polynomial (1,2 or 3)
ylab	Dependent variable name (Accepts the expression() function)
xlab	Independent variable name (Accepts the expression() function)
theme	ggplot2 theme (default is theme_classic())

118 polynomial2

se Adds confidence interval (*default* is FALSE)

point Defines whether to plot all points ("all"), mean ("mean"), mean with standard

deviation (default - "mean_sd") or mean with standard error ("mean_se").

legend.title Title legend

posi Legend position

textsize Font size (default is 12)

ylim y-axis scale

family Font family (*default* is sans)

width.bar width of the error bars of a regression graph.

pointsize Point size (default is 4)

linesize line size (Trendline and Error Bar)

separate Separation between treatment and equation (*default* is c("(\"","\")"))

n Number of decimal places for regression equations

DFres Residue freedom degrees

SSq Sum of squares of the residue

Value

Returns two or more linear, quadratic or cubic regression analyzes.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Leandro Simoes Azeredo Goncalves

Rodrigo Yudi Palhaci Marubayashi

See Also

polynomial, polynomial2_color

```
dose=rep(c(0,0,0,2,2,2,4,4,4,6,6,6),3)
resp=c(8,7,5,23,24,25,30,34,36,80,90,80,
12,14,15,23,24,25,50,54,56,80,90,40,
12,14,15,3,4,5,50,54,56,80,90,40)
trat=rep(c("A","B","C"),e=12)
polynomial2(dose, resp, trat, grau=c(1,2,3))
```

polynomial2_color 119

polynomial2_color

Analysis: Linear regression graph in double factorial with color graph

Description

Linear regression analysis for significant interaction of an experiment with two factors, one quantitative and one qualitative

Usage

```
polynomial2_color(
  fator1,
  resp,
  fator2,
  color = NA,
 grau = NA,
 ylab = "Response",
 xlab = "independent",
  theme = theme_classic(),
  se = FALSE,
 point = "mean_se",
 legend.title = "Treatments",
  posi = "top",
  textsize = 12,
 ylim = NA,
  family = "sans",
 width.bar = NA,
 pointsize = 5,
 linesize = 0.8,
  separate = c("(\"", "\")"),
 n = NA,
 DFres = NA,
  SSq = NA
)
```

Arguments

fator1	Numeric or complex vector with factor 1 levels
resp	Numerical vector containing the response of the experiment.
fator2	Numeric or complex vector with factor 2 levels
color	Graph color (default is NA)
grau	Degree of the polynomial (1,2 or 3)
ylab	Dependent variable name (Accepts the expression() function)
xlab	Independent variable name (Accepts the expression() function)
theme	ggplot2 theme (<i>default</i> is theme_classic())

polynomial2_color

se Adds confidence interval (*default* is FALSE)

point Defines whether to plot all points ("all"), mean ("mean"), mean with standard

deviation ("mean_sd") or mean with standard error (default - "mean_se").

legend.title Title legend

posi Legend position

textsize Font size (default is 12)

ylim y-axis scale

family Font family (*default* is sans)

width.bar width of the error bars of a regression graph.

pointsize Point size (default is 4)

linesize line size (Trendline and Error Bar)

separate Separation between treatment and equation (*default* is c("(\"","\")"))

n Number of decimal places for regression equations

DFres Residue freedom degrees

SSq Sum of squares of the residue

Value

Returns two or more linear, quadratic or cubic regression analyzes.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Leandro Simoes Azeredo Goncalves

Rodrigo Yudi Palhaci Marubayashi

See Also

polynomial, polynomial2

```
dose=rep(c(0,0,0,2,2,2,4,4,4,6,6,6),3)
resp=c(8,7,5,23,24,25,30,34,36,80,90,80,
12,14,15,23,24,25,50,54,56,80,90,40,
12,14,15,3,4,5,50,54,56,80,90,40)
trat=rep(c("A","B","C"),e=12)
polynomial2_color(dose, resp, trat, grau=c(1,2,3))
```

pomegranate 121

pomegranate

Dataset: Pomegranate data

Description

An experiment was conducted with the objective of studying different products to reduce the loss of mass in postharvest of pomegranate fruits. The experiment was conducted in a completely randomized design with four replications. Treatments are: T1: External Wax; T2: External + Internal Wax; T3: External Orange Oil; T4: Internal + External Orange Oil; T5: External sodium hypochlorite; T6: Internal + External sodium hypochlorite

Usage

data(pomegranate)

Format

data.frame containing data set

trat Categorical vector with treatments

WL Numeric vector weights loss

SS Numeric vector solid soluble

AT Numeric vector titratable acidity

ratio Numeric vector with ratio (SS/AT)

See Also

cloro, enxofre, laranja, mirtilo, porco, sensorial, simulate1, simulate2, simulate3, tomate, weather, phao, passiflora

Examples

data(pomegranate)

porco

Dataset: Pig development and production

Description

An experiment whose objective was to study the effect of castration age on the development and production of pigs, evaluating the weight of the piglets. Four treatments were studied: A - castration at 56 days of age; B - castration at 7 days of age; C - castration at 36 days of age; D - whole (not castrated); E - castration at 21 days of age. The Latin square design was used in order to control the variation between litters (lines) and the variation in the initial weight of the piglets (columns), with the experimental portion consisting of a piglet.

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Usage

```
data(porco)
```

Format

```
data.frame containing data set

trat Categorical vector with treatments
linhas Categorical vector with lines
colunas Categorical vector with columns
resp Numeric vector
```

See Also

cloro, enxofre, laranja, mirtilo, pomegranate, sensorial, simulate1, simulate2, simulate3, tomate, weather, phao, passiflora, aristolochia

Examples

```
data(porco)
```

PSUBDBC

Analysis: DBC experiments in split-plot

Description

Analysis of an experiment conducted in a randomized block design in a split-plot scheme using fixed effects analysis of variance.

Usage

```
PSUBDBC(
  f1,
  f2,
  block,
  response,
  norm = "sw",
  alpha.f = 0.05,
  alpha.t = 0.05,
  quali = c(TRUE, TRUE),
  names.fat = c("F1", "F2"),
 mcomp = "tukey",
  grau = c(NA, NA),
  grau12 = NA,
  grau21 = NA,
  transf = 1,
  constant = 0,
```

PSUBDBC 123

```
geom = "bar",
theme = theme_classic(),
ylab = "Response",
xlab = "",
xlab.factor = c("F1", "F2"),
color = "rainbow",
textsize = 12,
labelsize = 4,
dec = 3,
legend = "Legend",
errorbar = TRUE,
addmean = TRUE,
ylim = NA,
point = "mean_se",
fill = "lightblue",
angle = 0,
family = "sans",
posi = "right",
angle.label = 0
```

Arguments

f1		Numeric or complex vector with plot levels
f2		Numeric or complex vector with subplot levels
block		Numeric or complex vector with blocks
respons	е	Numeric vector with responses
norm		Error normality test (default is Shapiro-Wilk)
alpha.f		Level of significance of the F test (default is 0.05)
alpha.t		Significance level of the multiple comparison test (default is 0.05)
quali		Defines whether the factor is quantitative or qualitative (qualitative)
names.f	at	Name of factors
mcomp		Multiple comparison test (Tukey (default), LSD, Scott-Knott and Duncan)
grau		Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with three elements.
grau12		Polynomial degree in case of quantitative factor ($default$ is 1). Provide a vector with n levels of factor 2, in the case of interaction f1 x f2 and qualitative factor 2 and quantitative factor 1.
grau21		Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with n levels of factor 1, in the case of interaction f1 x f2 and qualitative factor 1 and quantitative factor 2.
transf		Applies data transformation (default is 1; for log consider 0)
constan	t	Add a constant for transformation (enter value)
geom		Graph type (columns or segments (For simple effect only))

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theme ggplot2 theme (default is theme_classic())

ylab Variable response name (Accepts the *expression*() function)

xlab Treatments name (Accepts the *expression*() function)

xlab.factor Provide a vector with two observations referring to the x-axis name of factors 1

and 2, respectively, when there is an isolated effect of the factors. This argument

uses 'parse'.

color When the columns are different colors (Set fill-in argument as "trat")

textsize Font size (default is 12) labelsize Font size (default is 4)

dec Number of cells (*default* is 3)

legend Legend title name

errorbar Plot the standard deviation bar on the graph (In the case of a segment and column

graph) - default is TRUE

addmean Plot the average value on the graph (*default* is TRUE)

ylim y-axis limit

point This function defines whether the point must have all points ("all"), mean ("mean"),

standard deviation (*default* - "mean_sd") or mean with standard error ("mean_se") if quali= FALSE. For quali=TRUE, 'mean_sd' and 'mean_se' change which in-

formation will be displayed in the error bar.

fill Defines chart color (to generate different colors for different treatments, define

fill = "trat")

angle x-axis scale text rotation
family Font family (default is sans)

posi Legend position angle.label Label angle

Value

The table of analysis of variance, the test of normality of errors (Shapiro-Wilk, Lilliefors, Anderson-Darling, Cramer-von Mises, Pearson and Shapiro-Francia), the test of homogeneity of variances (Bartlett), the test of multiple comparisons (Tukey, LSD, Scott-Knott or Duncan) or adjustment of regression models up to grade 3 polynomial, in the case of quantitative treatments. The column chart for qualitative treatments is also returned. The function also returns a standardized residual plot.

Note

The order of the chart follows the alphabetical pattern. Please use 'scale_x_discrete' from package ggplot2, 'limits' argument to reorder x-axis. The bars of the column and segment graphs are standard deviation.

In the final output when transformation (transf argument) is different from 1, the columns resp and respo in the mean test are returned, indicating transformed and non-transformed mean, respectively.

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Author(s)

Gabriel Danilo Shimizu

References

Principles and procedures of statistics a biometrical approach Steel, Torry and Dickey. Third Edition 1997

Multiple comparisons theory and methods. Departament of statistics the Ohio State University. USA, 1996. Jason C. Hsu. Chapman Hall/CRC.

Practical Nonparametrics Statistics. W.J. Conover, 1999

Ramalho M.A.P., Ferreira D.F., Oliveira A.C. 2000. Experimentacao em Genetica e Melhoramento de Plantas. Editora UFLA.

Scott R.J., Knott M. 1974. A cluster analysis method for grouping mans in the analysis of variance. Biometrics, 30, 507-512.

Examples

PSUBDIC

Analysis: DIC experiments in split-plot

Description

Analysis of an experiment conducted in a completely randomized design in a split-plot scheme using fixed effects analysis of variance.

Usage

```
PSUBDIC(
f1,
f2,
block,
response,
```

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```
norm = "sw",
  alpha.f = 0.05,
  alpha.t = 0.05,
  quali = c(TRUE, TRUE),
  names.fat = c("F1", "F2"),
 mcomp = "tukey",
 grau = c(NA, NA),
 grau12 = NA,
  grau21 = NA,
  transf = 1,
  constant = 0,
  geom = "bar",
  theme = theme_classic(),
 ylab = "Response",
  xlab = "",
  xlab.factor = c("F1", "F2"),
  fill = "lightblue",
  angle = 0,
  family = "sans",
  color = "rainbow",
  legend = "Legend",
  errorbar = TRUE,
  addmean = TRUE,
  textsize = 12,
  labelsize = 4,
  dec = 3,
 ylim = NA,
 posi = "right",
 point = "mean_se",
 angle.label = 0
)
```

Arguments f1

	- Annabar of the Free Anna
f2	Numeric or complex vector with subplot levels
block	Numeric or complex vector with blocks
response	Numeric vector with responses
norm	Error normality test (default is Shapiro-Wilk)
alpha.f	Level of significance of the F test (default is 0.05)
alpha.t	Significance level of the multiple comparison test (default is 0.05)
quali	Defines whether the factor is quantitative or qualitative (qualitative)
names.fat	Name of factors
mcomp	Multiple comparison test (Tukey (default), LSD, Scott-Knott and Duncan)
grau	Polynomial degree in case of quantitative factor (<i>default</i> is 1). Provide a vector with three elements.

Numeric or complex vector with plot levels

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grau12 Polynomial degree in case of quantitative factor (*default* is 1). Provide a vector

with n levels of factor 2, in the case of interaction f1 x f2 and qualitative factor

2 and quantitative factor 1.

grau21 Polynomial degree in case of quantitative factor (*default* is 1). Provide a vector

with n levels of factor 1, in the case of interaction f1 x f2 and qualitative factor

1 and quantitative factor 2.

transf Applies data transformation (default is 1; for log consider 0)

constant Add a constant for transformation (enter value)

geom Graph type (columns or segments (For simple effect only))

theme ggplot2 theme (*default* is theme_classic())

ylab Variable response name (Accepts the *expression*() function) xlab Treatments name (Accepts the *expression*() function)

xlab.factor Provide a vector with two observations referring to the x-axis name of factors 1

and 2, respectively, when there is an isolated effect of the factors. This argument

uses 'parse'.

fill Defines chart color (to generate different colors for different treatments, define

fill = "trat")

angle x-axis scale text rotation family Font family (default is sans)

color When the columns are different colors (Set fill-in argument as "trat")

legend Legend title name

errorbar Plot the standard deviation bar on the graph (In the case of a segment and column

graph) - default is TRUE

addmean Plot the average value on the graph (*default* is TRUE)

textsize Font size (default is 12)
labelsize Label size (default is 4)

dec Number of cells (default is 3)

ylim y-axis limit posi Legend position

point This function defines whether the point must have all points ("all"), mean ("mean"),

standard deviation (*default* - "mean_sd") or mean with standard error ("mean_se") if quali= FALSE. For quali=TRUE, 'mean_sd' and 'mean_se' change which in-

formation will be displayed in the error bar.

angle.label Label angle

Value

The table of analysis of variance, the test of normality of errors (Shapiro-Wilk, Lilliefors, Anderson-Darling, Cramer-von Mises, Pearson and Shapiro-Francia), the test of homogeneity of variances (Bartlett), the test of multiple comparisons (Tukey, LSD, Scott-Knott or Duncan) or adjustment of regression models up to grade 3 polynomial, in the case of quantitative treatments. The column chart for qualitative treatments is also returned. The function also returns a standardized residual plot.

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Note

The order of the chart follows the alphabetical pattern. Please use 'scale_x_discrete' from package ggplot2, 'limits' argument to reorder x-axis. The bars of the column and segment graphs are standard deviation.

In the final output when transformation (transf argument) is different from 1, the columns resp and respo in the mean test are returned, indicating transformed and non-transformed mean, respectively.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Leandro Simoes Azeredo Goncalves

Rodrigo Yudi Palhaci Marubayashi

References

Principles and procedures of statistics a biometrical approach Steel, Torry and Dickey. Third Edition 1997

Multiple comparisons theory and methods. Departament of statistics the Ohio State University. USA, 1996. Jason C. Hsu. Chapman Hall/CRC.

Practical Nonparametrics Statistics. W.J. Conover, 1999

Ramalho M.A.P., Ferreira D.F., Oliveira A.C. 2000. Experimentacao em Genetica e Melhoramento de Plantas. Editora UFLA.

Scott R.J., Knott M. 1974. A cluster analysis method for grouping mans in the analysis of variance. Biometrics, 30, 507-512.

Examples

PSUBFAT2DBC

Analysis: Plot subdivided into randomized blocks with a subplot in a double factorial scheme

Description

This function performs the analysis of a randomized block design in a split-plot with a subplot in a double factorial scheme.

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Usage

```
PSUBFAT2DBC(
    f1,
    f2,
    f3,
    block,
    resp,
    alpha.f = 0.05,
    alpha.t = 0.05,
    norm = "sw",
    homog = "bt",
    mcomp = "tukey"
)
```

Arguments

f1	Numeric or complex vector with plot levels
f2	Numeric or complex vector with splitplot levels
f3	Numeric or complex vector with splitsplitplot levels
block	Numeric or complex vector with blocks
resp	Numeric vector with responses
alpha.f	Level of significance of the F test (default is 0.05)
alpha.t	Significance level of the multiple comparison test ($default$ is 0.05)
norm	Error normality test (default is Shapiro-Wilk)
homog	Homogeneity test of variances (default is Bartlett)
mcomp	Multiple comparison test (Tukey (default), LSD and Duncan)

Value

Analysis of variance of fixed effects and multiple comparison test of Tukey, Scott-Knott, LSD or Duncan.

```
f1=rep(c("PD","PDE","C"), e = 40);f1=factor(f1,unique(f1))
f2=rep(c(300,400), e = 20,3);f2=factor(f2,unique(f2))
f3=rep(c("c1", "c2", "c3", "c4"), e = 5,6);f3=factor(f3,unique(f3))
bloco=rep(paste("B",1:5),24); bloco=factor(bloco,unique(bloco))
set.seed(10)
resp=rnorm(120,50,5)
PSUBFAT2DBC(f1,f2,f3,bloco,resp,alpha.f = 0.5) # force triple interaction
PSUBFAT2DBC(f1,f2,f3,bloco,resp,alpha.f = 0.4) # force double interaction
```

PSUBSUBDBC PSUBSUBDBC

PSUBSUBDBC

Analysis: DBC experiments in split-split-plot

Description

Analysis of an experiment conducted in a randomized block design in a split-split-plot scheme using analysis of variance of fixed effects.

Usage

```
PSUBSUBDBC(
f1,
f2,
f3,
block,
response,
alpha.f = 0.05,
alpha.t = 0.05,
dec = 3,
mcomp = "tukey"
)
```

Arguments

f1	Numeric or complex vector with plot levels
f2	Numeric or complex vector with splitplot levels
f3	Numeric or complex vector with splitsplitplot levels
block	Numeric or complex vector with blocks
response	Numeric vector with responses
alpha.f	Level of significance of the F test (default is 0.05)
alpha.t	Significance level of the multiple comparison test (<i>default</i> is 0.05)
dec	Number of cells
mcomp	Multiple comparison test (Tukey (default), LSD and Duncan)

Value

Analysis of variance of fixed effects and multiple comparison test of Tukey, LSD or Duncan.

Note

The PSUBSUBDBC function does not present residual analysis, interaction breakdown, graphs and implementations of various multiple comparison or regression tests. The function only returns the analysis of variance and multiple comparison test of Tukey, LSD or Duncan.

quant.fat2.desd 131

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

Examples

```
library(AgroR)
data(enxofre)
with(enxofre, PSUBSUBDBC(f1, f2, f3, bloco, resp))
```

quant.fat2.desd

Analysis: Polynomial splitting for double factorial in DIC and DBC

Description

Splitting in polynomials for double factorial in DIC and DBC. Note that f1 must always be qualitative and f2 must always be quantitative. This function is an easier way to visualize trends for dual factor schemes with a quantitative and a qualitative factor.

Usage

```
quant.fat2.desd(factors = list(f1, f2, block), response, dec = 3)
```

Arguments

factors Define f1 and f2 and/or block factors in list form. Please note that in the list it is

necessary to write 'f1', 'f2' and 'block'. See example.

response response variable dec Number of cells

Value

Returns the coefficients of the linear, quadratic and cubic models, the p-values of the t test for each coefficient (p.value.test) and the p-values for the linear, quadratic, cubic model splits and the regression deviations.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

See Also

FAT2DIC, FAT2DBC

132 radargraph

Examples

```
library(AgroR)
data(cloro)
quant.fat2.desd(factors = list(f1=cloro$f1,
f2=rep(c(1:4),e=5,2), block=cloro$bloco),
response=cloro$resp)
```

radargraph

Graph: Circular column chart

Description

Circular column chart of an experiment with a factor of interest or isolated effect of a factor

Usage

```
radargraph(model, ylim = NA, labelsize = 4, transf = FALSE)
```

Arguments

model DIC, DBC or DQL object

ylim y-axis limit

labelsize Font size of the labels

transf If the data has been transformed (*default* is FALSE)

Value

Returns pie chart with averages and letters from the Scott-Knott cluster test

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

See Also

```
barplot_positive, sk_graph, plot_TH, corgraph, spider_graph, line_plot
```

```
data("laranja")
a=with(laranja, DBC(trat,bloco,resp, mcomp = "sk"))
radargraph(a)
```

seg_graph 133

seg_	gr	apł	า

Graph: Point graph for one factor

Description

This is a function of the point graph for one factor

Usage

```
seg_graph(model, fill = "lightblue", horiz = TRUE, pointsize = 4.5)
```

Arguments

model DIC, DBC or DQL object

fill fill bars

horiz Horizontal Column (default is TRUE)

pointsize Point size

Value

Returns a point chart for one factor

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
```

Rodrigo Yudi Palhaci Marubayashi

See Also

```
radargraph, barplot_positive, plot_TH, corgraph, spider_graph, line_plot
```

seg_graph2

seg_graph2

Graph: Point graph for one factor model 2

Description

This is a function of the point graph for one factor

Usage

```
seg_graph2(
  model,
  theme = theme_gray(),
  pointsize = 4,
  pointshape = 16,
  horiz = TRUE,
  vjust = -0.6
)
```

Arguments

model DIC, DBC or DQL object

theme ggplot2 theme pointsize Point size

pointshape Format point (default is 16)

horiz Horizontal Column (default is TRUE)

vjust vertical adjusted

Value

Returns a point chart for one factor

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

See Also

radargraph, barplot_positive, plot_TH, corgraph, spider_graph, line_plot

sensorial 135

Examples

sensorial

Dataset: Sensorial data

Description

Set of data from a sensory analysis with six participants in which different combinations (blend) of the grape cultivar bordo and niagara were evaluated. Color (CR), aroma (AR), flavor (SB), body (CP) and global (GB) were evaluated. The data frame presents the averages of the evaluators.

Usage

```
data(sensorial)
```

Format

```
data.frame containing data set

Blend Categorical vector with treatment
variable Categorical vector with variables
resp Numeric vector
```

See Also

cloro, enxofre, laranja, mirtilo, pomegranate, porco, simulate1, simulate2, simulate3, tomate, weather, phao, passiflora, aristolochia

```
data(sensorial)
```

136 simulate2

simulate1

Dataset: Simulated data dict

Description

Simulated data from a completely randomized experiment with multiple assessments over time

Usage

```
data(simulate1)
```

Format

data.frame containing data set

tempo Categorical vector with time

trat Categorical vector with treatment

resp Categorical vector with response

See Also

cloro, enxofre, laranja, mirtilo, pomegranate, porco, sensorial, simulate2, simulate3, tomate, weather, phao, passiflora, aristolochia

Examples

```
data(simulate1)
```

simulate2

Dataset: Simulated data dbct

Description

Simulated data from a latin square experiment with multiple assessments over time

Usage

```
data(simulate2)
```

Format

data.frame containing data set

tempo Categorical vector with time

trat Categorical vector with treatment

bloco Categorical vector with block

resp Categorical vector with response

simulate3 137

See Also

cloro, enxofre, laranja, mirtilo, pomegranate, porco, sensorial, simulate1, simulate3, tomate, weather, phao, passiflora, aristolochia

Examples

```
data(simulate2)
```

simulate3

Dataset: Simulated data dqlt

Description

Simulated data from a completely randomized experiment with multiple assessments over time

Usage

```
data(simulate3)
```

Format

data.frame containing data set

tempo Categorical vector with time

trat Categorical vector with treatment

linhas Categorical vector with line

colunas Categorical vector with column

resp Categorical vector with response

See Also

cloro, enxofre, laranja, mirtilo, pomegranate, porco, sensorial, simulate1, simulate2, tomate, weather, phao, passiflora, aristolochia

```
data(simulate3)
```

138 sketch

sketch

Utils: Experimental sketch

Description

Experimental sketching function

Usage

```
sketch(
  trat,
  trat1 = NULL,
  trat2 = NULL,
  design = "DIC",
  pos = "line",
  color.sep = "all",
  ID = FALSE,
 print.ID = TRUE,
  add.streets.y = NA,
  add.streets.x = NA,
 label.x = "",
label.y = "",
  axissize = 12,
 legendsize = 12,
  labelsize = 4,
  export.csv = FALSE,
  comment.caption = NULL
)
```

Arguments trat

	7 - 2 - 2 - 2 - 2 - 2 - 2 - 2 - 2 - 2 -
trat1	Vector with levels of factor B (Set to NULL if not factorial or psub)
trat2	Vector with levels of factor C (Set to NULL if not factorial)
r	Number of repetitions
design	Experimental design (see note)
pos	Repeat position (line or column),
color.sep	Color box
ID	plot Add only identification in sketch
print.ID	Print table ID
add.streets.y	Adds streets by separating treatments in row or column. The user must supply a numeric vector grouping the rows or columns that must be together. See the example.

Vector with factor A levels

sketch 139

add.streets.x Adds streets by separating treatments in row or column. The user must supply a numeric vector grouping the rows or columns that must be together. See the example. label.x text in x label.y text in y axissize Axis size legendsize Title legend size labelsize Label size Save table template based on sketch in csv export.csv comment.caption

Add comment in caption

Value

Returns an experimental sketch according to the specified design.

Note

The sketches have only a rectangular shape, and the blocks (in the case of randomized blocks) can be in line or in a column.

For the design argument, you can choose from the following options:

```
design="DIC" Completely randomized design
design="DBC" Randomized block design
design="DQL" Latin square design
design="FAT2DIC" DIC experiments in double factorial
design="FAT2DBC" DBC experiments in double factorial
design="FAT3DIC" DIC experiments in triple factorial
design="FAT3DBC" DBC experiments in triple factorial
design="PSUBDIC" DIC experiments in split-plot
design="PSUBDBC" DBC experiments in split-plot
design="PSUBSUBDBC" DBC experiments in split-split
design="STRIP-PLOT" Strip-plot DBC experiments
```

For the color.sep argument, you can choose from the following options:

```
design="DIC" use "all" or "none"

design="DBC" use "all", "bloco" or "none"

design="DQL" use "all", "column", "line" or "none"

design="FAT2DIC" use "all", "f1", "f2" or "none"

design="FAT2DBC" use "all", "f1", "f2", "block" or "none"

design="FAT3DIC" use "all", "f1", "f2", "f3" or "none"

design="FAT3DBC" use "all", "f1", "f2", "f3", "block" or "none"

design="PSUBDIC" use "all", "f1", "f2" or "none"

design="PSUBDBC" use "all", "f1", "f2", "block" or "none"

design="PSUBDBC" use "all", "f1", "f2", "block" or "none"
```

sk_graph

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br> Leandro Simoes Azeredo Goncalves Rodrigo Yudi Palhaci Marubayashi

References

Mendiburu, F., & de Mendiburu, M. F. (2019). Package 'agricolae'. R Package, Version, 1-2.

```
Trat=paste("Tr",1:6)
# Completely randomized design
sketch(Trat,r=3)
sketch(Trat,r=3,pos="column")
sketch(Trat,r=3,color.sep="none")
sketch(Trat,r=3,color.sep="none",ID=TRUE,print.ID=TRUE)
sketch(Trat,r=3,pos="column",add.streets.x=c(1,1,2,2,3,3))
# Randomized block design
sketch(Trat, r=3, design="DBC")
sketch(Trat, r=3, design="DBC",pos="column")
sketch(Trat, r=3, design="DBC",pos="column",add.streets.x=c(1,1,2))
sketch(Trat, r=3, design="DBC",pos="column",add.streets.x=c(1,2,3), add.streets.y=1:6)
sketch(Trat, r=3, design="DBC",pos="line",add.streets.y=c(1,2,3), add.streets.x=1:6)
# Completely randomized experiments in double factorial
sketch(trat=c("A","B"),
      trat1=c("A", "B", "C"),
      design = "FAT2DIC",
      r=3)
sketch(trat=c("A","B"),
      trat1=c("A", "B", "C"),
      design = "FAT2DIC",
      r=3,
      pos="column")
```

soybean 141

Description

This is a function of the bar graph for the Scott-Knott test

Usage

```
sk_graph(model, horiz = TRUE, fill.label = "lightyellow")
```

Arguments

model DIC, DBC or DQL object

horiz Horizontal Column (default is TRUE)

fill.label fill Label box fill color

Value

Returns a bar chart with columns separated by color according to the Scott-Knott test

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

See Also

radargraph, barplot_positive, plot_TH, corgraph, spider_graph, line_plot

Examples

soybean

Dataset: Soybean

Description

An experiment was carried out to evaluate the grain yield (kg ha-1) of ten different commercial soybean cultivars in the municipality of Londrina/Parana. The experiment was carried out in the design of randomized complete blocks with four replicates per treatment.

spider_graph

Usage

```
data("soybean")
```

Format

```
data.frame containing data set

cult numeric vector with treatment

bloc numeric vector with block

prod Numeric vector with grain yield
```

See Also

cloro, laranja, enxofre, laranja, mirtilo, passiflora, phao, porco, pomegranate, simulate1, simulate2, simulate3, tomate, weather

Examples

```
data(soybean)
```

spider_graph

Graph: Spider graph for sensorial analysis

Description

Spider chart or radar chart. Usually used for graphical representation of acceptability in sensory tests

Usage

```
spider_graph(
  resp,
  vari,
  blend,
  legend.title = "",
  xlab = "",
  ylab = "",
  ymin = 0
)
```

Arguments

resp Vector containing notes

vari Vector containing the variables blend Vector containing treatments

legend.title Caption title

STRIPLOT 143

```
xlab x axis title
ylab y axis title
ymin Minimum value of y
```

Value

Returns a spider or radar chart. This graph is commonly used in studies of sensory analysis.

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

See Also

```
radargraph, sk_graph, plot_TH, corgraph, barplot_positive, line_plot
```

Examples

```
library(AgroR)
data(sensorial)
with(sensorial, spider_graph(resp, variable, Blend))
```

STRIPLOT

Analysis: DBC experiments in strip-plot

Description

Analysis of an experiment conducted in a block randomized design in a strit-plot scheme using fixed effects analysis of variance.

Usage

```
STRIPLOT(
f1,
f2,
block,
response,
norm = "sw",
alpha.f = 0.05,
transf = 1,
textsize = 12,
labelsize = 4,
constant = 0
)
```

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Arguments

f1	Numeric or complex vector with plot levels
f2	Numeric or complex vector with subplot levels

block Numeric or complex vector with blocks

response Numeric vector with responses

norm Error normality test (*default* is Shapiro-Wilk) alpha.f Level of significance of the F test (*default* is 0.05)

transf Applies data transformation (default is 1; for log consider 0)

textsize Font size (default is 12) labelsize Label size (default is 4)

constant Add a constant for transformation (enter value)

Value

The table of analysis of variance, the test of normality of errors (Shapiro-Wilk, Lilliefors, Anderson-Darling, Cramer-von Mises, Pearson and Shapiro-Francia), the test of homogeneity of variances (Bartlett). The function also returns a standardized residual plot.

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>

Leandro Simoes Azeredo Goncalves

Rodrigo Yudi Palhaci Marubayashi

References

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Multiple comparisons theory and methods. Departament of statistics the Ohio State University. USA, 1996. Jason C. Hsu. Chapman Hall/CRC.

Practical Nonparametrics Statistics. W.J. Conover, 1999

Ramalho M.A.P., Ferreira D.F., Oliveira A.C. 2000. Experimentacao em Genetica e Melhoramento de Plantas. Editora UFLA.

Scott R.J., Knott M. 1974. A cluster analysis method for grouping mans in the analysis of variance. Biometrics, 30, 507-512.

summarise_anova 145

summarise_anova	Utils: Summary of Analysis of Variance and Test of Means

Description

Summarizes the output of the analysis of variance and the multiple comparisons test for completely randomized (DIC), randomized block (DBC) and Latin square (DQL) designs.

Usage

```
summarise_anova(analysis, inf = "p", design = "DIC", round = 3, divisor = TRUE)
```

Arguments

analysis	List with the analysis outputs of the DIC, DBC, DQL, FAT2DIC, FAT2DBC, PSUBDIC and PSUBDBC functions
inf	Analysis of variance information (can be "p", "f", "QM" or "SQ")
design	Type of experimental project (DIC, DBC, DQL, FAT2DIC, FAT2DBC, PSUBDIC or PSUBDBC)
round	Number of decimal places
divisor	Add divider between columns

Value

returns a data.frame or print with a summary of the analysis of several experimental projects.

Note

Adding table divider can help to build tables in microsoft word. Copy console output, paste into MS Word, Insert, Table, Convert text to table, Separated text into:, Other: I.

The column names in the final output are imported from the ylab argument within each function.

This function is only for declared qualitative factors. In the case of a quantitative factor and the other qualitative in projects with two factors, this function will not work.

Triple factorials and split-split-plot do not work in this function.

Author(s)

Gabriel Danilo Shimizu

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Examples

```
library(AgroR)
# DIC
data(pomegranate)
attach(pomegranate)
a=DIC(trat, WL, geom = "point", ylab = "WL")
b=DIC(trat, SS, geom = "point", ylab="SS")
c=DIC(trat, AT, geom = "point", ylab = "AT")
summarise_anova(analysis = list(a,b,c), divisor = TRUE)
library(knitr)
kable(summarise_anova(analysis = list(a,b,c), divisor = FALSE))
vari=c("WL","SS","AT")
output=lapply(vari,function(x){
output=DIC(trat,response = unlist(pomegranate[,x]),ylab = parse(text=x))})
summarise_anova(analysis = output, divisor = TRUE)
# DBC
data(soybean)
attach(soybean)
a=DBC(cult,bloc,prod,ylab = "Yield")
summarise_anova(list(a),design = "DBC")
# FAT2DIC
data(corn)
attach(corn)
a=FAT2DIC(A, B, Resp, quali=c(TRUE, TRUE))
summarise_anova(list(a),design="FAT2DIC")
```

summarise_conj

Utils: Summary of Analysis of Variance and Test of Means for Joint analysis

Description

Summarizes the output of the analysis of variance and the multiple comparisons test for completely randomized (DIC) and randomized block (DBC) designs for Joint analysis with qualitative factor.

Usage

```
summarise_conj(analysis, design = "DBC", info = "p")
```

summarise_dunnett 147

Arguments

analysis List with the analysis outputs of the conjdic and conjdbc functions

design Type of experimental project (DIC or DBC)

info Analysis of variance information (can be "p", "f", "QM" or "SQ")

Note

The column names in the final output are imported from the ylab argument within each function.

This function is only for declared qualitative factors. In the case of a quantitative factor and the other qualitative in projects with two factors, this function will not work.

Author(s)

Gabriel Danilo Shimizu

Examples

```
library(AgroR)
data(mirtilo)
set.seed(1); resp1=rnorm(36,10,4)
set.seed(4); resp2=rnorm(36,10,3)
set.seed(8); resp3=rnorm(36,100,40)
type1=with(mirtilo, conjdbc(trat, bloco, exp, resp, ylab = "var1"))
type2=with(mirtilo, conjdbc(trat, bloco, exp, resp1, ylab = "var2"))
type3=with(mirtilo, conjdbc(trat, bloco, exp, resp2, ylab = "var3"))
type4=with(mirtilo, conjdbc(trat, bloco, exp, resp3, ylab = "var4"))
summarise_conj(analysis = list(type1,type2,type3,type4))
```

summarise_dunnett

Utils: Dunnett's Test Summary

Description

Performs a summary in table form from a list of Dunnett's test outputs

Usage

```
summarise_dunnett(variable, colnames = NA, info = "sig")
```

Arguments

variable List object Dunnett test colnames Names of column info Information of table

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Value

A summary table from Dunnett's test is returned

Examples

```
library(AgroR)
data("pomegranate")
a=with(pomegranate,dunnett(trat=trat,resp=WL,control="T1"))
b=with(pomegranate,dunnett(trat=trat,resp=SS,control="T1"))
c=with(pomegranate,dunnett(trat=trat,resp=AT,control="T1"))
d=with(pomegranate,dunnett(trat=trat,resp=ratio,control="T1"))
summarise_dunnett(list(a,b,c,d))
```

tabledesc

Descriptive: Table descritive analysis

Description

Function for generating a data.frame with averages or other descriptive measures grouped by a categorical variable

Usage

```
tabledesc(data, fun = mean)
```

Arguments

data frame containing the first column with the categorical variable and the re-

maining response columns

fun Function of descriptive statistics (default is mean)

Value

Returns a data.frame with a measure of dispersion or position from a dataset and separated by a factor

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

Examples

```
data(pomegranate)
tabledesc(pomegranate)
library(knitr)
kable(tabledesc(pomegranate))
```

TBARPLOT.reverse 149

TBARPLOT.reverse Graph

Graph: Reverse graph of DICT, DBCT and DQL output when geom="bar"

Description

The function performs the construction of a reverse graph on the output of DICT, DBCT and DQL when geom="bar".

Usage

```
TBARPLOT.reverse(plot.t)
```

Arguments

```
plot.t DICT, DBCT or DQLT output when geom="bar"
```

Value

Returns a reverse graph of the output of DICT, DBCT or DQLT when geom="bar".

Note

All layout and subtitles are imported from DICT, DBCT and DQLT functions

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

See Also

```
DICT, DBCT, DQLT
```

Examples

```
data(simulate1)
a=with(simulate1, DICT(trat, tempo, resp,geom="bar",sup=40))
TBARPLOT.reverse(a)
```

150 test_two

test_two

Analysis: Test for two samples

Description

Test for two samples (paired and unpaired t test, paired and unpaired Wilcoxon test)

Usage

```
test_two(
  trat,
 resp,
 paired = FALSE,
 correct = TRUE,
  test = "t",
 alternative = c("two.sided", "less", "greater"),
 conf.level = 0.95,
  theme = theme_classic(),
 ylab = "Response",
 xlab = "",
 var.equal = FALSE,
 pointsize = 2,
 yposition.p = NA,
 xposition.p = NA,
 fill = "white"
)
```

Arguments

trat	Categorical vector with the two treatments
resp	Numeric vector with the response
paired	A logical indicating whether you want a paired t-test.
correct	A logical indicating whether to apply continuity correction in the normal approximation for the p-value.
test	Test used (t for test t or w for Wilcoxon test)
alternative	A character string specifying the alternative hypothesis, must be one of "two.sided" (default), "greater" or "less". You can specify just the initial letter.
conf.level	Confidence level of the interval.
theme	ggplot2 theme (default is theme_classic())
ylab	Variable response name (Accepts the expression() function)
xlab	Treatments name (Accepts the expression() function)
var.equal	A logical variable indicating whether to treat the two variances as being equal. If TRUE then the pooled variance is used to estimate the variance otherwise the Welch (or Satterthwaite) approximation to the degrees of freedom is used.

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pointsize Point size

yposition.p Position p-value in y

xposition.p Position p-value in x

fill fill box

Details

Alternative = "greater" is the alternative that x has a larger mean than y. For the one-sample case: that the mean is positive.

If paired is TRUE then both x and y must be specified and they must be the same length. Missing values are silently removed (in pairs if paired is TRUE). If var.equal is TRUE then the pooled estimate of the variance is used. By default, if var.equal is FALSE then the variance is estimated separately for both groups and the Welch modification to the degrees of freedom is used.

If the input data are effectively constant (compared to the larger of the two means) an error is generated.

Value

Returns the test for two samples (paired or unpaired t test, paired or unpaired Wilcoxon test)

Author(s)

```
Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi
```

Examples

```
resp=rnorm(100,100,5)
trat=rep(c("A","B"),e=50)
test_two(trat,resp)
test_two(trat,resp,paired = TRUE)
```

tomate Dataset: Tomato data

Description

An experiment conducted in a randomized block design in a split plot scheme was developed in order to evaluate the efficiency of bacterial isolates in the development of tomato cultivars. The experiment counted a total of 24 trays; each block (in a total of four blocks), composed of 6 trays, in which each tray contained a treatment (6 isolates). Each tray was seeded with 4 different genotypes, each genotype occupying 28 cells per tray. The trays were randomized inside each block and the genotypes were randomized inside each tray.

152 tonetest

Usage

```
data(tomate)
```

Format

```
data.frame containing data set

parc Categorical vector with plot
subp Categorical vector with split-plot
bloco Categorical vector with block
resp Numeric vector
```

See Also

cloro, enxofre, laranja, mirtilo, pomegranate, porco, sensorial, simulate1, simulate2, simulate3, weather, aristolochia, phao, passiflora

Examples

data(tomate)

tonetest

Analysis: t test to compare means with a reference value

Description

Sometimes the researcher wants to test whether the treatment mean is greater than/equal to or less than a reference value. For example, I want to know if the average productivity of my treatment is higher than the average productivity of a given country. For this, this function allows comparing the means with a reference value using the t test.

Usage

```
tonetest(response, trat, mu = 0, alternative = "two.sided", conf.level = 0.95)
```

Arguments

response Numerical vector containing the response of the experiment.

Numerical or complex vector with treatmentsA number indicating the true value of the mean

alternative A character string specifying the alternative hypothesis, must be one of "two.sided"

(default), "greater" or "less"

conf.level confidence level of the interval.

transf 153

Value

returns a list with the mean per treatment, maximum, minimum, sample standard deviation, confidence interval, t-test statistic and its p-value.

Note

No treatment can have zero variability. Otherwise the function will result in an error.

Author(s)

Gabriel Danilo Shimizu

Examples

```
library(AgroR)
data("pomegranate")
tonetest(resp=pomegranate$WL,
trat=pomegranate$trat,
mu=2,
alternative = "greater")
```

transf

Utils: Data transformation (Box-Cox, 1964)

Description

Estimates the lambda value for data transformation

Usage

```
transf(response, f1, f2 = NA, f3 = NA, block = NA, line = NA, column = NA)
```

Arguments

response	Numerical vector containing the response of the experiment.
f1	Numeric or complex vector with factor 1 levels
f2	Numeric or complex vector with factor 2 levels
f3	Numeric or complex vector with factor 3 levels
block	Numerical or complex vector with blocks
line	Numerical or complex vector with lines
column	Numerical or complex vector with columns

Value

Returns the value of lambda and/or data transformation approximation, according to Box-Cox (1964)

154 weather

Author(s)

Gabriel Danilo Shimizu, <shimizu@uel.br>
Leandro Simoes Azeredo Goncalves
Rodrigo Yudi Palhaci Marubayashi

References

Box, G. E., Cox, D. R. (1964). An analysis of transformations. Journal of the Royal Statistical Society: Series B (Methodological), 26(2), 211-243.

Examples

```
# Completely randomized design
data("pomegranate")
with(pomegranate, transf(WL,f1=trat))
# Randomized block design
data(soybean)
with(soybean, transf(prod, f1=cult, block=bloc))
# Completely randomized design in double factorial
data(cloro)
with(cloro, transf(resp, f1=f1, f2=f2))
# Randomized block design in double factorial
data(cloro)
with(cloro, transf(resp, f1=f1, f2=f2, block=bloco))
```

weather Dataset: Weather data

Description

Climatic data from 01 November 2019 to 30 June 2020 in the municipality of Londrina-PR, Brazil. Data from the Instituto de Desenvolvimento Rural do Parana (IDR-PR)

Usage

data(weather)

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Format

data.frame containing data set

Data POSIXct vector with dates

tempo Numeric vector with time

Tmax Numeric vector with maximum temperature

Tmed Numeric vector with mean temperature

Tmin Numeric vector with minimum temperature

UR Numeric vector with relative humidity

See Also

cloro, enxofre, laranja, mirtilo, pomegranate, porco, sensorial, simulate1, simulate2, simulate3, tomate, aristolochia, phao, passiflora

Examples

data(weather)

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