Package 'GerminaR'

October 12, 2022

```
Type Package
Title Indices and Graphics for Assess Seed Germination Process
Version 2.1.4
Description
      A collection of different indices and visualization techniques for evaluate the seed germina-
      tion process in ecophysiological studies (Lozano-Isla et al. 2019) <doi:10.1111/1440-1703.1275>.
Date 2022-05-18
URL https://germinar.inkaverse.com/,
      https://github.com/flavjack/germinar
BugReports https://github.com/flavjack/germinar/issues
Depends shiny, agricolae, ggplot2, dplyr, R (>= 3.5.0)
Imports tidyr, tibble, purrr, DT
Suggests knitr, rmarkdown, gsheet, inti
VignetteBuilder knitr
License GPL-3 | file LICENSE
LazyData true
Encoding UTF-8
RoxygenNote 7.2.0
NeedsCompilation no
Author Flavio Lozano-Isla [aut, cre] (<a href="https://orcid.org/0000-0002-0714-669X">https://orcid.org/0000-0002-0714-669X</a>),
      Omar Benites Alfaro [aut] (<a href="https://orcid.org/0000-0002-6852-9598">https://orcid.org/0000-0002-6852-9598</a>),
      Marcelo F. Pompelli [aut, ths]
       (<https://orcid.org/0000-0002-4408-6374>),
      Denise Garcia de Santana [aut],
      Marli A. Ranal [aut],
      Federal University of Pernambuco [cph] (Register of application),
      Federal Rural University of Pernambuco [cph] (Post Graduate Program),
      Inkaverse [ctb]
Maintainer Flavio Lozano-Isla <flavjack@gmail.com>
Repository CRAN
Date/Publication 2022-05-18 22:10:10 UTC
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fplot

Plot line or bar graphic

Description

Function use the dtsm function for plot the results

Usage

```
fplot(
  data,
  type = "bar",
  x,
  y,
  group = NA,
  xlab = NA,
  ylab = NA,
  glab = NA,
  ylimits = NA,
  xrotation = NA,
```

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```
xtext = NA,
gtext = NA,
legend = "top",
sig = NA,
sigsize = 3,
error = NA,
color = TRUE,
opt = NA
)
```

Arguments

data	Output from ger_testcomp function
type	Type of graphic. "bar" or "line"
X	Axis x variable
У	Axis y variable
group	Group variable
xlab	Title for the axis x
ylab	Title for the axis y
glab	Title for the legend
ylimits	limits of the y axis
xrotation	Rotation in x axis c(angle, h, v)
xtext	Text labels in x axis
gtext	Text labels in groups
legend	the position of legends ("none", "left", "right", "bottom", "top", or two-element numeric vector)
sig	Column with the significance
sigsize	Font size in significance letters
error	Show the error bar ("ste" or "std").
color	colored figure c(TRUE, FALSE) or vector with the color.
opt	Add news layer to the plot

Value

Line o bar plot

```
## Not run:
library(GerminaR)
library(dplyr)
```

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```
smr <- ger_summary(SeedN = "seeds"</pre>
                    , evalName = "D"
                    , data = prosopis) %>%
  mutate(across(rep:temp, as.factor))
av <- aov(grp ~ nacl*temp, smr)</pre>
anova(av)
mc <- ger_testcomp(aov = av</pre>
                    , comp = c("nacl", "temp"))
plotdt <- mc$table</pre>
plot <- fplot(data = plotdt</pre>
       , type = "bar"
       , x = "temp"
       , y = "grp"
       , group = "nacl"
       , sig = "sig"
       #, error = "ste"
       , color = T
       , ylimits = c(0, 120, 20)
plot
## End(Not run)
```

GerminaQuant

GerminaQuant for R

Description

GermiQuant for R app allows make the calculation for the germination indices incredibly easy in a interactive applications build in base a GerminaR R package and Shiny. GermiQuant app is live!. Outputs change instantly as users modify inputs, without requiring a reload the app. The principal features of the application allow calculate the principal germination indices, statistical analysis and easy way to plot the results.

Usage

```
GerminaQuant(dependencies = FALSE)
```

Arguments

dependencies Install package dependencies for run the app

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Source

Lozano-Isla, F., Benites Alfaro, O., & Pompelli, M. F. (2016). GerminaQuant for R (Patent No. BR 51 2016 001327-3). https://flavjack.shinyapps.io/germinaquant/

ger_ASG

ArcSin of Germination Percentage

Description

This function calculates the arcsin of germination percentage for normalization.

Usage

```
ger_ASG(SeedN, evalName, data)
```

Arguments

SeedN Name of the column with the number of seeds sown.

evalName Prefix of the names of the periods of evaluation.

data The name of the data frame containing the data.

Value

It returns an vector with the ArcSin of Germination values

Examples

```
library(GerminaR)
dfr <- prosopis
gas <- ger_ASG(SeedN = "seeds", evalName = "D", data = dfr)
gas</pre>
```

ger_boxp

Boxplot graphic

Description

Function use the raw data for made a boxplot graphic

ger_boxp

Usage

```
ger_boxp(
  data,
  x,
  y,
  group = NULL,
  xlab = NULL,
  ylab = NULL,
  glab = NULL,
  ylimits = NULL,
  xrotation = NULL,
  legend = "top",
  xtext = NULL,
  gtext = NULL,
  opt = NULL
)
```

Arguments

data	raw data
x	Axis x variable
у	Axis y variable
group	Group variable
xlab	Title for the axis x
ylab	Title for the axis y
glab	Title for the legend
ylimits	Limitis and break of the y axis c(init, end, brakes)
xrotation	Rotation in x axis c(angle, h, v)
legend	the position of legends ("none", "left", "right", "bottom", "top", or two-element numeric vector)
xtext	Text labels in x axis
gtext	Text labels in groups
opt	Add news layer to the plot

Value

boxplot

```
## Not run:
library(GerminaR)
```

ger_CVG

ger_CVG

Coefficient of Variance of the Mean Germination Time

Description

This function calculates the coefficient of variation of the mean germination time

Usage

```
ger_CVG(evalName, data)
```

Arguments

evalName Prefix of the names of the periods of evaluation.

data The name of the data frame containing the data.

Value

It returns an vector with the values of Coefficient of Variance of germination

```
library(GerminaR)
dfr <- prosopis
cvg <- ger_CVG(evalName = "D", data = dfr)
cvg</pre>
```

ger_GRP

~~~	CDD
ger	GRP

Germination Seed Percentage

### Description

This function calculates the germination percentage related at total seed sown for experimental unit.

### Usage

```
ger_GRP(SeedN, evalName, data)
```

### **Arguments**

SeedN Name of the column with the number of seeds sown.

evalName Prefix of the names of the periods of evaluation.

data The name of the data frame containing the data.

#### **Details**

According GOUVEA LABOURIAU (1983), the germinability of a sample of is the percentage of seeds in which the seed germination process comes to an end, in experimental conditions by the seminal intrauterine growth resulting protrusion (or emergence) of a living embryo.

#### Value

It returns an vector with the percentage of seed germinated.

### References

LABOURIAU, L. G.; VALADARES, M. E. B. The germination of seeds. OEA, Washington, DC, 1983.

```
library(GerminaR)
dt <- prosopis
grp <- ger_GRP(SeedN = "seeds",evalName = "D", data = dt)
grp</pre>
```

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ger_GRS

Germinated Seed Number

### **Description**

This function calculates the number of seed germinated.

### Usage

```
ger_GRS(evalName, data)
```

### **Arguments**

evalName Prefix of the names of the periods of evaluation.

data The name of the data frame containing the data.

### Value

Number of seed germinated

### **Examples**

```
library(GerminaR)
dt <- prosopis
grs <- ger_GRS(evalName = "D", data = dt)
grs</pre>
```

ger_GSP

Germination Speed

### Description

This function calculates the Germination Speed according at the time lapse of the evaluations.

### Usage

```
ger_GSP(evalName, data)
```

### **Arguments**

evalName Prefix of the names of the periods of evaluation.

data The name of the data frame containing the data.

### Value

It returns an vector with the Germination Speed

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### **Examples**

```
library(GerminaR)
dfr <- prosopis
gsp <- ger_GSP(evalName = "D", data = dfr)
gsp</pre>
```

ger_intime

Cumulative sum of germination by period of time for line graphic

### **Description**

This function makes a data table with the cumulative sum of values of germination by days.

### Usage

```
ger_intime(Factor, SeedN, evalName, method = "percentage", data)
```

### **Arguments**

Factor which will be graph in time

SeedN Name of the column with the seed numbers

evalName Prefix of the evaluation variable

method Type of cumulative germination. "percentage" or "relative"

data Data with the germination evaluation process

#### **Details**

Need a summary by factor before use it with function SummaryBy.

#### Value

Data frame with the germination by period

ger_MGR

ger_MGR

Mean Germination Rate

#### **Description**

This function calculates the mean germination rate of the germination.

### Usage

```
ger_MGR(evalName, data)
```

### **Arguments**

evalName Prefix of the names of the periods of evaluation.

data The name of the data frame containing the data.

#### **Details**

The average speed of germination is defined as the reciprocal of the average time germination (RANAL; SANTANA, 2006).

#### Value

It returns an vector with the values of Mean Germination Rate

#### References

RANAL, M. A.; SANTANA, D. G. DE. How and why to measure the germination process? Revista Brasileira de Botanica, v. 29, n. 1, p. 1-11, mar. 2006.

```
library(GerminaR)
dfr <- prosopis
mgr <- ger_MGR(evalName = "D", data = dfr)
mgr</pre>
```

ger_MGT

ger_MGT

Mean Germination Time

### Description

This function calculates the mean germination time of germination according at the time lapse of the evaluations.

### Usage

```
ger_MGT(evalName, data)
```

### **Arguments**

evalName Prefix of the names of the periods of evaluation.

data The name of the data frame containing the data.

#### **Details**

It was proposed by Haberlandt in 1875. It is calculated as the weighted average germination time. The number of germinated seeds at the intervals established for the collection of data is used as weight. It is expressed in terms of the same units of time used in the germination count (CZABATOR, 1962).

#### Value

It returns an vector with the values of Mean Germination Time.

#### References

CZABATOR, F. J. Germination value: an index combining speed and completeness of pine seed germination. Forest Science, v. 8, n. 4, p. 386-396, 1962.

```
library(GerminaR)
dfr <- prosopis
mgt <- ger_MGT(evalName = "D", data = dfr)
mgt</pre>
```

ger_SDG

ger_SDG

Standard deviation of the Mean Germination Time

### Description

This function calculates the standard deviation of the mean germination time

### Usage

```
ger_SDG(evalName, data)
```

### Arguments

evalName Prefix of the names of the periods of evaluation.

data The name of the data frame containing the data.

#### Value

It returns an vector with the values of Standard deviation of germination

### **Examples**

```
library(GerminaR)
dfr <- prosopis
sdg <- ger_SDG(evalName = "D", data = dfr)
sdg</pre>
```

ger_summary

Summary of Germination indices

### Description

This function makes a data table with the result of germination indices for each experimental unit.

### Usage

```
ger_summary(SeedN, evalName, data)
```

#### **Arguments**

SeedN Name of the column with the seed numbers

evalName Prefix of the evaluation variable

data The name of the data frame containing the data.

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#### Value

Data frame with the summary values of germination variables.

### **Examples**

```
library(GerminaR)
fb <- prosopis
smr <- ger_summary(SeedN = "seeds", evalName = "D", data = fb)
smr</pre>
```

ger_SYN

Germination Synchronization Index

### Description

This function calculates the germination synchronization of the germination process.

#### Usage

```
ger_SYN(evalName, data)
```

### **Arguments**

evalName Prefix of the names of the periods of evaluation.

data The name of the data frame containing the data.

#### **Details**

The Synchory Index Z has been proposed to assess the degree of overlap between flowering individuals in a population. By adopting the idea expressed by PRIMACK, R.B. (1980) the synchrony of one seed with other included in the same replication. Z = 1 when germination of all the seeds occurs at the same time and Z = 0 when at least two seeds can germinate one each time. Z produces a number if and only if there are two seeds finishing the seed germination process at the same time. Thus, the value of Z assessments is the grade of overlap between seed germination.

### Value

It returns an vector with the values of Germination synchrony

#### References

RANAL, M. A.; SANTANA, D. G. DE. How and why to measure the germination process? Revista Brasileira de Botanica, v. 29, n. 1, p. 1-11, mar. 2006.

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#### **Examples**

```
library(GerminaR)
dfr <- prosopis
syn <- ger_SYN(evalName = "D", data = dfr)
syn</pre>
```

ger_testcomp

Multiple comparison test

### **Description**

Function analysis of variance for summary data.

#### Usage

```
ger_testcomp(aov, comp, type = "snk", sig = 0.05)
```

### **Arguments**

aov lm o aov result function.

comp treatments will be compared.

type method for made comparison analysis: c("snk", "tukey", "duncan").

sig significance level. Default 0.05

#### Value

Table with complete data for graphics

ger_UNC

ger_UNC

Germination Uncertainty

#### **Description**

This function calculates the germination uncertainty in the germination process.

#### Usage

```
ger_UNC(evalName, data)
```

### Arguments

evalName Prefix of the names of the periods of evaluation.

data The name of the data frame containing the data.

### **Details**

The uncertainty index u is an adaptation of Shannon index measures the degree of uncertainty in predicting the informational entropy or uncertainty associated with the distribution of the relative frequency of germination (GOUVEA LABOURIAU 1983; LABOURIAU; VALADARES, 1983). Low values of u indicate frequencies with short peaks, i.e. the more concentrated the germination in time. Just a germinated seed changes the value of u. This means that u measures the degree of germination scattering.

#### Value

It returns an vector with the values of Germination Uncertainty.

#### References

GOUVEA LABOURIAU, L. L. G. L. A germinacao das sementes. Washington. LABOURIAU, L. G.; VALADARES, M. E. B. The germination of seeds. OEA, Washington, DC, 1983.

```
library(GerminaR)
dfr <- prosopis
unc <- ger_UNC(evalName = "D", data = dfr)
unc</pre>
```

ger_VGT

ger_VGT

Variance of the Mean Germination Time

### Description

This function calculates the variance of the mean germination time.

### Usage

```
ger_VGT(evalName, data)
```

### **Arguments**

evalName Prefix of the names of the periods of evaluation.

data The name of the data frame containing the data.

#### Value

It returns an vector with the values of Variance of Germination

### **Examples**

```
library(GerminaR)
dfr <- prosopis
vgt <- ger_VGT(evalName = "D", data = dfr)
vgt</pre>
```

gquant_analysis

GerminaQuant data analysis

### **Description**

Function analysis of variance for summary data.

### Usage

```
gquant_analysis(
  data,
  response,
  factors,
  block = NA,
  comparison = NA,
  type = "snk",
  sig = 0.05
)
```

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### **Arguments**

data from ger_summary() function
response germination indices to analyse
factors factor as vector or factor model as string
block block factor for RCBD
comparison treatments will be compared.
type method for made comparison analysis: c("snk", "tukey", "duncan").

sig significance level. Default 0.05

#### **Details**

Function for the analysis module in the app

#### Value

list

### **Examples**

osmp

Osmotic potential calculator

### Description

Function to calculate the grams of salt or PEG-6000 needed for determinated osmotic potential

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#### Usage

```
osmp(type = "salt", vol, pres, temp, mw, ki)
```

#### **Arguments**

type Salt or PEG-6000 c("salt", "peg6000"). Default: "salt".

vol volume (liters)

pres Pressure (Mpa) in negative values. 1 bar = 0.1 Mpa

temp Temperature (centigrade)

mw Molecular weight

ki Salt dissociation constant (NaCl = 1.8)

#### Value

Numeric value (grams)

prosopis

Germination under different osmotic potentials and temperatures.

### **Description**

Dataset containing information from germination experiment with Prosopis juliflor under different osmotic potentials and temperatures.

- rep a numeric vector, repetitions or replications.
- nacl a numeric vector, NaCl osmotic potentials levels in Mpa to be evaluated.
- temp a numeric vector, temperature levels in centigrades to be evelauted.
- seeds a numeric vector, number of seed used for experimental unit.
- D0-D10 numeric vectors, ten evaluations days performed in the germination experiment.

#### Usage

prosopis

#### Format

A data frame with 15 columns and 80 rows.

#### **Source**

LEV-UFRPE

#### References

MIRANDA, R. D. Q.; CORREIA, R. M.; DE ALMEIDA-CORTEZ, J. S.; POMPELLI, M. F. Germination of Prosopis juliflora (Sw.) D.C. seeds at different osmotic potentials and temperatures. Plant Species Biology, v. 29, n. 3, p. E9-E20, set. 2014.

20 textcolor

rep_row

Repeated Rows in a data matrix

### Description

This function made a data table with the evaluation days of germination

### Usage

```
rep_row(Rseq, Nrow)
```

### **Arguments**

Rseq Row sequence for the data matrix

Nrow Number of rows for the data matrix

#### Value

Data Matrix with day of the germination

textcolor

Colourise text for display in the terminal.

### Description

If R is not currently running in a system that supports terminal colours the text will be returned unchanged.

### Usage

```
textcolor(text, fg = "red", bg = NULL)
```

#### **Arguments**

text character vector

fg foreground colour, defaults to white
bg background colour, defaults to transparent

#### **Details**

Allowed colours are: black, blue, brown, cyan, dark gray, green, light blue, light cyan, light gray, light green, light purple, light red, purple, red, white, yellow

### Author(s)

testthat package

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### **Examples**

```
print(textcolor("Red", "red"))
cat(textcolor("Red", "red"), "\n")
cat(textcolor("White on red", "white", "red"), "\n")
```

webTable

Web table

### Description

Web table

### Usage

```
webTable(
  data,
  caption = NULL,
  digits = 3,
  rnames = FALSE,
  buttons = NULL,
  file_name = NULL,
  scrolly = NULL
)
```

### Arguments

data	Dataset.
caption	Title for the table.
digits	Digits number in the table exported.
rnames	Row names.
buttons	Buttons: "excel", "copy" or "none". Default c("excel", "copy")
file_name	Excel file name
scrolly	Windows height to show the table

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