# Package 'inti'

September 3, 2024

```
Type Package
Version 0.6.6
Title Tools and Statistical Procedures in Plant Science
Description The 'inti' package is part of the 'inkaverse' project for developing
      different procedures and tools used in plant science and experimental designs.
      The mean aim of the package is to support researchers during the planning of
      experiments and data collection (tarpuy()), data analysis and graphics (yupana())
      , and technical writing.
      Learn more about the 'inkaverse' project at <a href="https://inkaverse.com/">https://inkaverse.com/>.
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colortext 3

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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

```
colortext(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

#### **Examples**

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

design\_noreps

Experimental design without replications

#### **Description**

Function to deploy field-book experiment without replications

design\_noreps

#### Usage

```
design_noreps(
  factors,
  type = "sorted",
  zigzag = FALSE,
  nrows = NA,
  serie = 100,
  seed = NULL,
  fbname = "inkaverse",
  qrcode = "{fbname}{plots}{factors}"
)
```

## Arguments

factors	Lists with names and factor vector [list].
type	Randomization in the list [string: sorted, unsorted]
zigzag	Experiment layout in zigzag [logic: FALSE].
nrows	Experimental design dimension by rows [numeric: value]
serie	Number to start the plot id [numeric: 1000].
seed	Replicability from randomization [numeric: NULL].
fbname	Bar code prefix for data collection [string: "inkaverse"].
qrcode	[string: "{fbname}{plots}{factors}"] String to concatenate the qr code.

#### Value

A list with the field-book design and parameters

design\_repblock 5

```
## End(Not run)
```

design\_repblock

Experimental design in CRD and RCBD

## Description

Function to deploy field-book experiment for CRD and RCBD

## Usage

```
design_repblock(
  nfactors = 1,
  factors,
  type = "crd",
  rep = 3,
  zigzag = FALSE,
  nrows = NA,
  serie = 100,
  seed = NULL,
  fbname = "inkaverse",
  qrcode = "{fbname}{plots}{factors}"
)
```

## Arguments

nfactors	Number of factor in the experiment [numeric: 1].
factors	Lists with names and factor vector [list].
type	Type of experimental arrange [string: "crd" "rcbd" "lsd"]
rep	Number of replications in the experiment [numeric: 3].
zigzag	Experiment layout in zigzag [logic: F].
nrows	Experimental design dimension by rows [numeric: value]
serie	Number to start the plot id [numeric: 100].
seed	Replicability from randomization [numeric: NULL].
fbname	Bar code prefix for data collection [string: "inkaverse"].
qrcode	[string: " $\{fbname\}\{plots\}\{factors\}$ "] String to concatenate the qr code.

#### Value

A list with the field-book design and parameters

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#### **Examples**

```
## Not run:
library(inti)
factores <- list("geno" = c("A", "B", "C", "D", "D", 1, NA, NA, NULL, "NA")</pre>
                 , "salt stress" = c(0, 50, 200, 200, "T0", NA, NULL, "NULL")
                 , time = c(30, 60, 90)
fb <-design_repblock(nfactors = 2</pre>
                      , factors = factores
                      , type = "rcbd"
                      , rep = 5
                      , zigzag = T
                      , seed = 0
                      , nrows = 20
                      , qrcode = "{fbname}{plots}{factors}"
dsg <- fb$fieldbook</pre>
fb %>%
  tarpuy_plotdesign(fill = "plots")
fb$parameters
## End(Not run)
```

figure2qmd

Figure to Quarto format

## Description

Use Articul8 Add-ons from Google docs to build Rticles

## Usage

```
figure2qmd(text, path = ".", opts = NA)
```

## Arguments

text	Markdown text with figure information [string]
path	Image path for figures [path: "." (base directory)]
opts	chunk options in brackets [string: NA]

figure2rmd 7

#### **Details**

Quarto option can be included in the title using "{{}}" separated by commas

#### Value

string mutated

figure2rmd

Figure to Rmarkdown format

#### **Description**

Use Articul8 Add-ons from Google docs to build Rticles

#### Usage

```
figure2rmd(text, path = ".", opts = NA)
```

## Arguments

text String with the table information
path Path of the image for the figure
opts chunk options in brackets.

#### Value

Mutated string

footnotes

Footnotes in tables

## **Description**

Include tables footnotes and symbols for kables in pandoc format

#### Usage

```
footnotes(table, notes = NULL, label = "Note:", notation = "alphabet")
```

## **Arguments**

table Kable output in pandoc format.

notes Footnotes for the table.

label Label for start the footnote.

notation Notation for the footnotes (default = "alphabet"). See details.

8 gdoc2qmd

#### **Details**

You should use the pandoc format kable(format = "pipe"). You can add the footnote symbol using {hypen} in your table. notation could be use: "alphabet", "number", "symbol", "none".

#### Value

Table with footnotes for word and html documents

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20	IUCZUII	ıu

Google docs to Rmarkdown

## Description

Use Articul8 Add-ons from Google docs to build Rticles

#### Usage

```
gdoc2qmd(file, export = NA, format = "qmd", type = "asis")
```

## Arguments

file Zi	p file path from	n Articul8 exported	d in md format	[path]

export Path to export the files [path: NA (file directory)]

format Output format [string: "qmd" "rmd"]

type output file type [strig: "asis" "list", "listfull", "full"]

## **Details**

Document rendering until certain point: "#| end" Include for next page: "#| newpage" You can include the cover page params using "#|" in a Google docs table

#### Value

path

H2cal 9

H2cal

Broad-sense heritability in plant breeding

## Description

Heritability in plant breeding on a genotype difference basis

## Usage

```
H2cal(
  data,
  trait,
  gen.name,
  rep.n,
  env.n = 1,
  year.n = 1,
  env.name = NULL,
  year.name = NULL,
  fixed.model,
  random.model,
  summary = FALSE,
  emmeans = FALSE,
  weights = NULL,
  plot_diag = FALSE,
  outliers.rm = FALSE,
  trial = NULL
)
```

## Arguments

data	Experimental design data frame with the factors and traits.
trait	Name of the trait.
gen.name	Name of the genotypes.
rep.n	Number of replications in the experiment.
env.n	Number of environments (default = $1$ ). See details.
year.n	Number of years (default = 1). See details.
env.name	Name of the environments (default = NULL). See details.
year.name	Name of the years (default = NULL). See details.
fixed.mode	The fixed effects in the model (BLUEs). See examples.
random.mod	del The random effects in the model (BLUPs). See examples.
summary	Print summary from random model (default = FALSE).
emmeans	Use emmeans for calculate the BLUEs (default = FALSE).
weights	an optional vector of 'prior weights' to be used in the fitting process (default = NULL).

10 H2cal

plot\_diag Show diagnostic plots for fixed and random effects (default = FALSE). Options:

"base", "ggplot". .

outliers.rm Remove outliers (default = FALSE). See references.

trial Column with the name of the trial in the results (default = NULL).

#### **Details**

The function allows to made the calculation for individual or multi-environmental trials (MET) using fixed and random model.

1. The variance components based in the random model and the population summary information based in the fixed model (BLUEs).

- 2. Heritability under three approaches: Standard (ANOVA), Cullis (BLUPs) and Piepho (BLUEs).
- 3. Best Linear Unbiased Estimators (BLUEs), fixed effect.
- 4. Best Linear Unbiased Predictors (BLUPs), random effect.
- 5. Table with the outliers removed for each model.

For individual experiments is necessary provide the trait, gen.name, rep.n.

For MET experiments you should env.n and env.name and/or year.n and year.name according your experiment.

The BLUEs calculation based in the pairwise comparison could be time consuming with the increase of the number of the genotypes. You can specify emmeans = FALSE and the calculate of the BLUEs will be faster.

If emmeans = FALSE you should change 1 by 0 in the fixed model for exclude the intersect in the analysis and get all the genotypes BLUEs.

For more information review the references.

#### Value

list

#### Author(s)

Maria Belen Kistner

Flavio Lozano Isla

#### References

Bernal Vasquez, Angela Maria, et al. "Outlier Detection Methods for Generalized Lattices: A Case Study on the Transition from ANOVA to REML." Theoretical and Applied Genetics, vol. 129, no. 4, Apr. 2016.

Buntaran, H., Piepho, H., Schmidt, P., Ryden, J., Halling, M., and Forkman, J. (2020). Cross validation of stagewise mixed model analysis of Swedish variety trials with winter wheat and spring barley. Crop Science, 60(5).

Schmidt, P., J. Hartung, J. Bennewitz, and H.P. Piepho. 2019. Heritability in Plant Breeding on a Genotype Difference Basis. Genetics 212(4).

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Schmidt, P., J. Hartung, J. Rath, and H.P. Piepho. 2019. Estimating Broad Sense Heritability with Unbalanced Data from Agricultural Cultivar Trials. Crop Science 59(2).

Tanaka, E., and Hui, F. K. C. (2019). Symbolic Formulae for Linear Mixed Models. In H. Nguyen (Ed.), Statistics and Data Science. Springer.

Zystro, J., Colley, M., and Dawson, J. (2018). Alternative Experimental Designs for Plant Breeding. In Plant Breeding Reviews. John Wiley and Sons, Ltd.

#### **Examples**

include\_pdf

Include PDF in markdown documents

## Description

Insert PDF files in markdown documents

#### Usage

```
include_pdf(file, width = "100%", height = "600")
```

#### **Arguments**

```
file file path from pdf file.
width width preview file.
height height preview file.
```

include\_table

#### Value

html code for markdown

include\_table

Table with footnotes

#### **Description**

Include tables with title and footnotes for word and html documents

#### Usage

```
include_table(table, caption = NA, notes = NA, label = NA, notation = "none")
```

#### **Arguments**

```
table Data frame.

caption Table caption (default = NULL). See details.

notes Footnotes for the table (default = NA). See details.

label Label for start the footnote (default = NA).

Notation for the symbols and footnotes (default = "none") Others: "alphabet", "number", "symbol".
```

#### Value

Table with caption and footnotes

```
library(inti)

table <- data.frame(
x = rep_len(1, 5)
, y = rep_len(3, 5)
, z = rep_len("c", 5)
)

table %>% inti::include_table(
  caption = "Title caption b) line 0
a) line 1
b) line 2"
, notes = "Footnote"
, label = "Where:"
)
```

jc\_tombola 13

 $jc\_tombola$ 

Journal Club Tombola

#### **Description**

Function for arrange journal club schedule

## Usage

```
jc_tombola(
  data,
  members,
  papers = 1,
  group = NA,
  gr_lvl = NA,
  status = NA,
  st_lvl = "active",
  frq = 7,
  date = NA,
  seed = NA
)
```

#### **Arguments**

data	Data frame withe members and their information.
members	Columns with the members names.
papers	Number of paper by meeting
group	Column for arrange the group.
gr_lvl	Levels in the groups for the arrange. See details.
status	Column with the status of the members.
st_lvl	Level to confirm the assistance in the JC. See details.
frq	Number of the day for each session.
date	Date when start the first session of JC.
seed	Number for replicate the results (default = date).

#### **Details**

The function could consider n levels for gr\_lvl. In the case of more levels using "both" or "all" will be the combination. The suggested levels for st\_lvl are: active or spectator. Only the "active" members will enter in the schedule.

## Value

data frame with the schedule for the JC

14 mean\_comparison

mean\_comparison

Mean comparison test

#### **Description**

Function to compare treatment from lm or aov using data frames

## Usage

```
mean_comparison(
  data,
  response,
  model_factors,
  comparison,
  test_comp = "SNK",
  sig_level = 0.05
)
```

## Arguments

data Fieldbook data.

response Model used for the experimental design.

model\_factors Factor in the model.

comparison Significance level for the analysis (default = 0.05).

test\_comp Comparison test (default = "SNK"). Others: "TUKEY", "DUNCAN".

sig\_level Significance level for the analysis (default = 0.05).

#### Value

list

met 15

```
, model_factors = "bloque* geno*treat"
, comparison = c("geno", "treat")
, test_comp = "SNK"
)
mc$comparison
mc$stat

## End(Not run)
```

met

Swedish cultivar trial data

#### **Description**

The datasets were obtained from official Swedish cultivar tests. Dry matter yield was analyzed. All trials were laid out as alpha-designs with two replicates. Within each replicate, there were five to seven incomplete blocks.

#### Usage

met

#### **Format**

A data frame with 1069 rows and 8 variables:

```
zone Sweden is divided into three different agricultural zones: South, Middle, and North location Locations: 18 location in the Zones
rep Replications (4): number of replication in the experiment
alpha Incomplete blocks (8) in the alpha-designs
cultivar Cultivars (30): genotypes evaluated
yield Yield in kg/ha
year Year (1): 2016
env environment (18): combination zone + location + year
```

#### Source

doi:10.1002/csc2.20177

outliers\_remove

metamorphosis	Transform fieldbooks based in a dictionary	

#### **Description**

Transform entire fieldbook according to data a dictionary

#### Usage

```
metamorphosis(fieldbook, dictionary, from, to, index, colnames)
```

#### **Arguments**

fieldbook Data frame with the original information.

dictionary Data frame with new names and categories. See details. from Column of the dictionary with the original names.

to Column of the dictionary with the new names.

index Column of the dictionary with the type and level of the variables.

colnames Character vector with the name of the columns.

## **Details**

The function require at least three columns.

- 1. Original names (from).
- 2. New names (to).
- 3. Variable type (index).

#### Value

List with two objects. 1. New data frame. 2. Dictionary.

|--|--|--|

#### **Description**

Use the method M4 in Bernal Vasquez (2016). Bonferroni Holm test to judge residuals standardized by the re scaled MAD (BH MADR).

#### Usage

```
outliers_remove(data, trait, model, drop_na = TRUE)
```

plot\_diag

#### **Arguments**

data	Experimental design data frame with the factors and traits.

trait Name of the trait.

model The fixed or random effects in the model.
drop\_na drop NA values from the data.frame

#### **Details**

Function to remove outliers in MET experiments

#### Value

list. 1. Table with date without outliers. 2. The outliers in the dataset.

#### References

Bernal Vasquez, Angela Maria, et al. "Outlier Detection Methods for Generalized Lattices: A Case Study on the Transition from ANOVA to REML." Theoretical and Applied Genetics, vol. 129, no. 4, Apr. 2016.

#### **Examples**

```
library(inti)

rmout <- potato %>% outliers_remove(
  data = .
  , trait ="stemdw"
  , model = "0 + treat*geno + (1|bloque)"
  , drop_na = FALSE
  )

rmout
```

plot\_diag

Diagnostic plots

## Description

Function to plot the diagnostic of models

#### Usage

```
plot_diag(model, title = NA)
```

plot\_diagnostic

#### **Arguments**

model Statistical model title Plot title

#### Value

plots

## **Examples**

```
## Not run:
library(inti)
lm <- aov(stemdw ~ bloque + geno*treat, data = potato)
# lm <- potato %>% lme4::lmer(stemdw ~ (1|bloque) + geno*treat, data = .)
plot(lm, which = 1)
plot_diag(lm)[3]

plot(lm, which = 2)
plot_diag(lm)[2]

plot(lm, which = 3)
plot_diag(lm)[4]

plot(lm, which = 4)
plot_diag(lm)[1]

## End(Not run)
```

plot\_diagnostic

Diagnostic plots

#### **Description**

Function to plot the diagnostic of models

#### Usage

```
plot_diagnostic(data, formula, title = NA)
```

## Arguments

data Experimental design data frame with the factors and traits.

formula Mixed model formula

title Plot title

plot\_raw 19

## Value

plots

## **Examples**

plot\_raw

Plot raw data

## Description

Function use the raw data for made a boxplot graphic

## Usage

```
plot_raw(
  data,
  type = "boxplot",
 у,
  group = NULL,
 xlab = NULL,
 ylab = NULL,
 glab = NULL,
 ylimits = NULL,
 xlimits = NULL,
  xrotation = NULL,
  legend = "top",
  xtext = NULL,
  gtext = NULL,
  color = TRUE,
 linetype = 1,
  opt = NULL
)
```

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## Arguments

data	raw data
type	Type of graphic. "boxplot" or "scatterplot"
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 and break of the y axis c(initial, end, brakes)
xlimits	For scatter plot. Limits and break of the x axis c(initial, 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 using a vector
gtext	Text labels in groups using a vector
color	Colored figure (TRUE), black & white (FALSE) or color vector
linetype	Line type for regression. Default = $0$
opt	Add new layers to the plot

## Details

You could add additional layer to the plot using "+" with ggplot2 options

## Value

plot

plot\_smr 21

plot\_smr

Plot summary data

#### **Description**

Graph summary data into bar o line plot

#### Usage

```
plot_smr(
  data,
  type = NULL,
  x = NULL,
  y = NULL,
  group = NULL,
 xlab = NULL,
 ylab = NULL,
  glab = NULL,
 ylimits = NULL,
  xrotation = c(0, 0.5, 0.5),
  xtext = NULL,
  gtext = NULL,
  legend = "top",
  sig = NULL,
  sigsize = 3,
  error = NULL,
  color = TRUE,
  opt = NULL
)
```

#### **Arguments**

```
data

Output from summary data

type

Type of graphic. "bar" or "line"

x

Axis x variable
```

plot\_smr

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 c(initial, end, brakes)
xrotation	Rotation in x axis c(angle, h, v)
xtext	Text labels in x axis using a vector
gtext	Text labels in group using a vector
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 (TRUE), black & white (FALSE) or color vector
opt	Add news layer to the plot

#### **Details**

If the table is a out put of mean\_comparison(graph\_opts = TRUE) function. Its contain all the parameter for the plot.

You could add additional layer to the plot using "+" with ggplot2 options

## Value

plot

potato 23

```
, group = "treat"
, glab = "Tratamientos"
, ylimits = c(0, 1, 0.2)
, color = c("red", "black")
, gtext = c("Irrigado", "Sequia")
)

## End(Not run)
```

potato

Water use efficiency in 15 potato genotypes

## Description

Experiment to evaluate the physiological response from 15 potatos genotypes under water deficit condition. The experiment had a randomized complete block design with five replications. The stress started at 30 day after planting.

#### Usage

potato

#### **Format**

```
A data frame with 150 rows and 17 variables:
```

```
treat Water deficit treatments: sequia, irrigado
geno 15 potato genotypes
bloque blocks for the experimentl design
spad_29 Relative chlorophyll content (SPAD) at 29 day after planting
spad_83 Relative chlorophyll content (SPAD) at 84 day after planting
rwc_84 Relative water content (percentage) at 84 day after planting
op_84 Osmotic potential (Mpa) at 84 day after planting
leafdw leaf dry weight (g)
stemdw stem dry weight (g)
rootdw root dry weight (g)
tubdw tuber dry weight (g)
biomdw total biomass dry weight (g)
hi harvest index
ttrans total transpiration (1)
wue water use effiency (g/l)
twue tuber water use effiency (g/l)
lfa leaf area (cm2)
```

24 remove\_outliers

ramova	outliers	
remove	outiters	

Remove outliers using mixed models

#### **Description**

Use the method M4 in Bernal Vasquez (2016). Bonferroni Holm test to judge residuals standardized by the re scaled MAD (BH MADR).

## Usage

```
remove_outliers(data, formula, drop_na = FALSE, plot_diag = FALSE)
```

#### **Arguments**

data Experimental design data frame with the factors and traits.

formula mixed model formula.

drop\_na drop NA values from the data.frame

plot\_diag Diagnostic plot based in the raw and clean data

#### **Details**

Function to remove outliers in MET experiments

#### Value

list. 1. Table with date without outliers. 2. The outliers in the dataset.

#### References

Bernal Vasquez, Angela Maria, et al. "Outlier Detection Methods for Generalized Lattices: A Case Study on the Transition from ANOVA to REML." Theoretical and Applied Genetics, vol. 129, no. 4, Apr. 2016.

```
library(inti)

rmout <- potato %>%
    remove_outliers(data = .
    , formula = stemdw ~ 0 + (1|bloque) + treat*geno
    , plot_diag = FALSE
    , drop_na = FALSE
)
```

split\_folder 25

split\_folder

Split folder

## Description

Function to split folder by size or number of elements

## Usage

```
split_folder(
  folder,
  export,
  units = "megas",
  size = 500,
  zip = TRUE,
  remove = FALSE
)
```

#### **Arguments**

```
folder Path of folder to split (path).

export Path to export the split folders (path).

units Units to split folder (string: "megas", "number").

size Folder size by the units selected (numeric).

zip Zip split folders (logical).

remove Remove the split folder after zip (logical).
```

#### Value

zip files

```
## Not run:
split_folder("pictures/QUINOA 2018-2019 SC SEEDS EDWIN - CAMACANI/"
    , "pictures/split_num", remove = T, size = 400, units = "number")
## End(Not run)
```

26 table2rmd

table2qmd

Table to Quarto format

## Description

Use Articul8 Add-ons from Google docs to build Rticles

## Usage

```
table2qmd(text, type = "asis")
```

## Arguments

text Markdown text with table information (string)

type output file type [strig: "asis" "list", "listfull", "full"]

#### Value

string mutated

table2rmd

Table to Rmarkdown format

## Description

Use Articul8 Add-ons from Google docs to build Rticles

#### Usage

```
table2rmd(text, opts = NA)
```

## Arguments

text String with the table information

opts chunk options in brackets.

## Value

Mutated string

tarpuy 27

tarpuy

Interactive fieldbook designs

## Description

Invoke RStudio addin to create fieldbook designs

## Usage

```
tarpuy(dependencies = FALSE)
```

## Arguments

dependencies Install package dependencies for run the app

#### **Details**

Tarpuy allow to create experimental designs under an interactive app.

#### Value

Shiny app

## Examples

```
if(interactive()){
  inti::tarpuy()
}
```

tarpuy\_design

Fieldbook experimental designs

## Description

Function to deploy experimental designs

28 tarpuy\_design

#### Usage

```
tarpuy_design(
  data,
  nfactors = 1,
  type = "crd",
  rep = 2,
  zigzag = FALSE,
  nrows = NA,
  serie = 100,
  seed = NULL,
  fbname = NA,
  qrcode = "{fbname}{plots}{factors}"
)
```

#### **Arguments**

data Exp	perimental design	data frame wit	th the factors an	d level. See examples.
----------	-------------------	----------------	-------------------	------------------------

nfactors Number of factor in the experiment(default = 1). See details.

type Type of experimental arrange (default = "crd"). See details.

Number of replications in the experiment (default = 3).

zigzag Experiment layout in zigzag [logic: FALSE].

nrows Experimental design dimension by rows [numeric: value]

serie Number to start the plot id [numeric: 100].

seed Replicability of draw results (default = 0) always random. See details.

fbname Barcode prefix for data collection.

qrcode [string: "{fbname}{plots}{factors}"] String to concatenate the qr code.

#### **Details**

The function allows to include the arguments in the sheet that have the information of the design. You should include 2 columns in the sheet: {arguments} and {values}. See examples. The information will be extracted automatically and deploy the design. nfactors = 1: crd, rcbd, lsd, lattice. nfactors = 2 (factorial): split-crd, split-rcbd split-lsd nfactors >= 2 (factorial): crd, rcbd, lsd.

#### Value

A list with the fieldbook design

```
## Not run:
library(inti)
library(gsheet)
```

tarpuy\_plex 29

tarpuy\_plex

Fieldbook plan information

#### **Description**

Information for build a plan for an experiment (PLEX)

#### Usage

```
tarpuy_plex(
 data = NULL,
  idea = NULL,
  goal = NULL,
 hypothesis = NULL,
  rationale = NULL,
 objectives = NULL,
 plan = NULL,
  institutions = NULL,
  researchers = NULL,
 manager = NULL,
 location = NULL,
  altitude = NULL,
  georeferencing = NULL,
  environment = NULL,
  start = NA,
  end = NA,
  about = NULL,
  fieldbook = NULL,
  gdocs = NULL,
  github = NULL,
  album = NULL,
 nfactor = 2,
  design = "rcbd",
  rep = 3,
```

30 tarpuy\_plex

```
zigzag = FALSE,
  nrows = NA,
  serie = 100.
  seed = 0,
  qrcode = "{fbname}{plots}{factors}"
)
```

#### **Arguments**

hypothesis

Data with the fieldbook information. data

idea How the idea was born. goal The main goal of the project.

What are the expected results. rationale Based in which evidence is planned the experiment.

objectives The objectives of the project.

plan General description of the project (M & M).

institutions Institutions involved in the project. Persons involved in the project. researchers

manager Persons responsible of the collection of the data.

location Location of the project.

altitude Altitude of the experiment (m.a.s.l).

georeferencing Georeferencing information.

environment Environment of the experiment (greenhouse, lab, etc).

The date of the start of the experiments. start The date of the end of the experiments. end

about Short description of the project.

fieldbook Name or ID for the fieldbook/project.

link for Google Docs gdocs

github link with the github repository. album link with the photos of the project. Number of factors for the design. nfactor

Type of design. design

Number of replication. rep

Experiment layout in zigzag [logic: F] zigzag

Experimental design dimension by rows [numeric: value] nrows

Number of digits in the plots. serie Seed for the randomization. seed

[string: "{fbname}{plots}{factors}"] String to concatenate the qr code. qrcode

tarpuy\_plotdesign 31

#### **Details**

Provide the information available.

#### Value

data frame or list of arguments:

- 1. info
- 2. variables
- 3. design
- 4. logbook
- 5. timetable
- 6. budget

tarpuy\_plotdesign

Fieldbook plot experimental designs

## Description

Plot fieldbook sketch designs based in experimental design

## Usage

```
tarpuy_plotdesign(
  data,
  factor = NA,
  fill = "plots",
  xlab = NULL,
  ylab = NULL,
  glab = NULL
)
```

## Arguments

data	Experimental design data frame with the factors and level. See examples.
factor	Vector with the name of the columns with the factors.
fill	Value for fill the experimental units (default = "plots").
xlab	Title for x axis.
ylab	Title for y axis.
glab	Title for group axis.

#### **Details**

The function allows to plot the experimental design according the field experiment design.

32 tarpuy\_traits

#### Value

plot

#### **Examples**

tarpuy\_traits

Field book traits

## Description

Function to export field book and traits for be used in field book app.

#### Usage

```
tarpuy_traits(fieldbook = NULL, last_factor = NULL, traits = NULL)
```

## Arguments

fieldbook Experiment field book [dataframe].

last\_factor Last factor in the field book [string: colnames]

traits Traits information [dataframe or list].

#### **Details**

For the traits parameters you can used shown in the Field Book app

tarpuy\_traits 33

#### Value

list

```
library(inti)
fieldbook <- inti::potato</pre>
traits <- list(</pre>
  list(variable = "altura de planta"
      , trait = "altp"
       , format = "numeric"
       , when = "30, 40, 50"
       , samples = 3
       , units = "cm"
       , details = NA
       , minimum = 0
       , maximum = 100
  , list(variable = "severidad"
        , trait = "svr"
         , format = "scategorical"
         , when = "30, 40, 50"
         , samples = 1
         , units = "scale"
         , details = NA
         , categories = "1, 3, 5, 7, 9"
  , list(variable = "foto"
          , trait = "foto"
          , format = "photo"
          , when = "hrv, pshrv"
          , samples = 1
          , units = "image"
          , details = NA
  )
    list(variable = "germinacion"
          , trait = "ger"
          , format = "boolean"
          , when = "30, 40, 50"
          , samples = 1
          , units = "logical"
          , details = NA
 )
fbapp <- tarpuy_traits(fieldbook, last_factor = "bloque", traits)</pre>
## Not run:
library(inti)
```

34 web\_table

web\_table

HTML tables for markdown documents

#### **Description**

Export tables with download, pasta and copy buttons

#### Usage

```
web_table(
  data,
  caption = NULL,
  digits = 2,
  rnames = FALSE,
  buttons = NULL,
  file_name = "file",
  scrolly = NULL,
  columnwidth = "200px",
  width = "100%"
)
```

yupana 35

#### **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. Default "45vh"

columnwidth Column width. Default '200px'

width Width in pixels or percentage (Defaults to automatic sizing)

#### Value

table in markdown format for html documents

## **Examples**

```
## Not run:
library(inti)
met %>%
  web_table(caption = "Web table")
## End(Not run)
```

yupana

Interactive data analysis

#### **Description**

Invoke RStudio addin to analyze and graph experimental design data

## Usage

```
yupana(dependencies = FALSE)
```

#### **Arguments**

dependencies Install package dependencies for run the app

#### **Details**

Yupana: data analysis and graphics for experimental designs.

36 yupana\_analysis

#### Value

Shiny app

#### **Examples**

```
if(interactive()){
  inti::yupana()
}
```

yupana\_analysis

Fieldbook analysis report

## Description

Function to create a complete report of the fieldbook

## Usage

```
yupana_analysis(
  data,
  last_factor = NULL,
  response,
  model_factors,
  comparison,
  test_comp = "SNK",
  sig_level = 0.05,
  plot_dist = "boxplot",
  plot_diag = FALSE,
  digits = 2
)
```

#### **Arguments**

data

The last factor in your fieldbook. last\_factor response Response variable. model\_factors Model used for the experimental design. comparison Factors to compare Comprasison test c("SNK", "TUKEY", "DUNCAN") test\_comp sig\_level Significal test (default: p = 0.005) Plot data distribution (default = "boxplot") plot\_dist plot\_diag Diagnostic plots for model (default = FALSE). digits Digits number in the table exported.

Field book data.

yupana\_export 37

#### Value

list

## **Examples**

yupana\_export

Graph options to export

## Description

Function to export the graph options and model parameters

## Usage

```
yupana_export(
  data,
  type = NA,
  xlab = NA,
 ylab = NA,
  glab = NA,
 ylimits = NA,
  xrotation = c(0, 0.5, 0.5),
 xtext = NA,
  gtext = NA,
  legend = "top",
  sig = NA,
  error = NA,
  color = TRUE,
 opt = NA,
  dimension = c(20, 10, 100)
)
```

38 yupana\_export

#### **Arguments**

data	Result from yupana_analysis or yupana_import.
type	Plot type
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 group
legend	the position of legends ("none", "left", "right", "bottom", "top", or two-element numeric vector)
sig	Column with the significance
error	Show the error bar ("ste" or "std").
color	colored figure (TRUE), otherwise black & white (FALSE)
opt	Add news layer to the plot
dimension	Dimension of graphs

#### Value

data frame

yupana\_import 39

yupana\_import

Import information from data summary

## Description

Graph summary data

#### Usage

```
yupana_import(data)
```

#### **Arguments**

data

Summary information with options

#### Value

list

40 yupana\_mvr

```
info <- yupana_import(fb)
## End(Not run)</pre>
```

yupana\_mvr

Multivariate Analysis

#### **Description**

Multivariate analysis for PCA and HCPC

## Usage

```
yupana_mvr(
  data,
  last_factor = NULL,
  summary_by = NULL,
  groups = NULL,
  variables = NULL
)
```

## **Arguments**

data Field book data.

last\_factor The last factor in your fieldbook [string: NULL].

summary\_by Variables for group the analysis.

groups Groups for color in PCA.

variables Variables to be use in the analysis [string: NULL].

#### **Details**

Compute and plot information for multivariate analysis (PCA, HCPC and correlation).

#### Value

result and plots

```
## Not run:
library(inti)
fb <- inti::potato</pre>
```

yupana\_reshape 41

yupana\_reshape

Fieldbook reshape

## Description

Function to reshape fieldbook according a separation character

## Usage

```
yupana_reshape(
  data,
  last_factor,
  sep,
  new_colname,
  from_var = NULL,
  to_var = NULL,
  exc_factors = NULL)
```

## Arguments

data	Field book raw data.
last_factor	The last factor in your field book.
sep	Character that separates the last value.
new_colname	The new name for the column created.
from_var	The first variable in case you want to exclude several. variables.
to_var	The last variable in case you want to exclude several variables.
exc_factors	Factor to exclude during the reshape.

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## **Details**

If you variable name is variable\_evaluation\_rep. The reshape function will help to create the column rep and the new variable name will be variable\_evaluation.

## Value

data frame

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