Package 'metan'

June 4, 2021

```
Type Package
Title Multi Environment Trials Analysis
Version 1.14.0
Maintainer Tiago Olivoto <tiagoolivoto@gmail.com>
Description Performs stability analysis of multi-environment
      trial data using parametric and non-parametric methods. Parametric
      methods includes Additive Main Effects and Multiplicative Interaction
      (AMMI) analysis by Gauch (2013) <doi:10.2135/cropsci2013.04.0241>,
      Ecovalence by Wricke (1965), Genotype plus Genotype-Environment (GGE)
      biplot analysis by Yan & Kang (2003) <doi:10.1201/9781420040371>,
      geometric adaptability index by Mohammadi & Amri (2008)
      <doi:10.1007/s10681-007-9600-6>, joint regression analysis by Eberhart
      & Russel (1966) <doi:10.2135/cropsci1966.0011183X000600010011x>,
      genotypic confidence index by Annicchiarico (1992), Murakami & Cruz's
      (2004) method, power law residuals
      (POLAR) statistics by Doring et al. (2015)
      <doi:10.1016/j.fcr.2015.08.005>, scale-adjusted coefficient of
      variation by Doring & Reckling (2018) <doi:10.1016/j.eja.2018.06.007>,
      stability variance by Shukla (1972) <doi:10.1038/hdy.1972.87>,
      weighted average of absolute scores by Olivoto et al. (2019a)
      <doi:10.2134/agronj2019.03.0220>, and multi-trait stability index by
      Olivoto et al. (2019b) <doi:10.2134/agronj2019.03.0221>.
      Non-parametric methods includes superiority index by Lin & Binns
      (1988) <doi:10.4141/cjps88-018>, nonparametric measures of phenotypic
      stability by Huehn (1990)
      <https://link.springer.com/article/10.1007/BF00024241>, TOP third
      statistic by Fox et al. (1990) <doi:10.1007/BF00040364>. Functions for
      computing biometrical analysis such as path analysis, canonical
      correlation, partial correlation, clustering analysis, and tools for
      inspecting, manipulating, summarizing and plotting typical
      multi-environment trial data are also provided.
License GPL-3
URL https://github.com/TiagoOlivoto/metan
BugReports https://github.com/TiagoOlivoto/metan/issues
Depends R (>= 3.5.0)
Imports dplyr (>= 1.0.0),
      GGally,
```

2 R topics documented:

```
ggforce,
     ggplot2 (>= 3.3.0),
     ggrepel,
     lme4,
     lmerTest,
     magrittr,
     mathjaxr,
     methods,
     patchwork,
     purrr,
     rlang (>= 0.1.2),
     tibble,
     tidyr,
     tidyselect (>= 1.0.0)
Suggests DT,
     knitr,
     rmarkdown,
     roxygen2
VignetteBuilder knitr
RdMacros mathjaxr
Encoding UTF-8
Language en-US
LazyData true
RoxygenNote 7.1.1
Roxygen list(markdown = TRUE)
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metan-package

Multi-Environment Trial Analysis

Description

metan provides functions for performing the most used analyses in the evaluation of multi-environment trials, including, but not limited to:

- · ANOVA-based stability statistics;
- · AMMI-based stability indexes;
- BLUP-based stability indexes;
- · Cross-validation procedures for AMMI-family and BLUP models;
- GGE biplot analysis;
- Estimation using AMMI considering different numbers of interaction principal component axes;
- Graphics tools for generating biplots;
- Nonparametric stability statistics;
- Variance components and genetic parameters in mixed-effect models;
- Within-environment analysis of variance;

metan also provides functions for biometrical analysis such as path analysis, canonical correlation, partial correlation, clustering analysis, as well as tools for summarizing and plotting data.

A complete guide may be found at https://tiagoolivoto.github.io/metan/

acv

Adjusted Coefficient of Variation

Description

[Stable]

Computes the scale-adjusted coefficient of variation, acv, (Doring and Reckling, 2018) to account for the systematic dependence of σ^2 from μ . The acv is computed as follows:

$$acv = \frac{\sqrt{10^{\tilde{v}_i}}}{\mu_i} \times 100$$

where \tilde{v}_i is the adjusted logarithm of the variance computed as:

$$\tilde{v}_i = a + (b-2)\frac{1}{n}\sum_i m_i + 2m_i + e_i$$

being a and b the coefficients of the linear regression for log_{10} of the variance over the log_{10} of the mean; m_i is the log_{10} of the mean, and e_i is the Power Law Residuals (POLAR), i.e., the residuals for the previously described regression.

acv 7

Usage

```
acv(mean, var, na.rm = FALSE)
```

Arguments

mean A numeric vector with mean values.

var A numeric vector with variance values.

na.rm If FALSE, the default, missing values are removed with a warning. If TRUE, missing values are silently removed.

Value

A tibble with the following columns

- mean The mean values;
- var The variance values;
- log10_mean The base 10 logarithm of mean;
- log10_var The base 10 logarithm of variance;
- **POLAR** The Power Law Residuals;
- cv The standard coefficient of variation;
- acv Adjusted coefficient of variation.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Doring, T.F., and M. Reckling. 2018. Detecting global trends of cereal yield stability by adjusting the coefficient of variation. Eur. J. Agron. 99: 30-36. doi: 10.1016/j.eja.2018.06.007

AMMI_indexes

AMMI-based stability indexes

Description

[Stable]

This function computes the following AMMI-based stability indexes: ASV, AMMI stability value (Purchase et al., 2000); SIPC, sums of the absolute value of the IPCA scores (Sneller et al. 1997); EV, averages of the squared eigenvector values (Sneller et al. 1997); and Za, absolute value of the relative contribution of IPCAs to the interaction (Zali et al. 2012), and WAAS, weighted average of absolute scores (Olivoto et al. 2019).

The ASV index is computed as follows:

$$ASV_i = \left[\left[\frac{r \, \lambda_1^2}{r \, \lambda_2^2} \times \left(\lambda_1^{0.5} \, a_{i1} t_{j1} \right) \right]^2 + \left(\lambda_2^{0.5} \, a_{i2} t_{j2} \right)^2 \right]^{0.5}$$

where r is the number of replications included in the analysis,

The SIPC index is computed as follows:

$$SIPC_i = \sum_{k=1}^{P} \left| |\lambda_k^{0.5} a_{ik}| \right|$$

where P is the number of IPCA retained via F-tests.

The EV index is computed as follows:

$$EV_i = \sum\nolimits_{k=1}^P a_{ik}^2 / P$$

The ZA index is computed as follows:

$$Za_i = \sum_{k=1}^{P} \theta_k a_{ik}$$

where θ_k is the percentage sum of squares explained by the kth IPCA.

$$WAAS_i = \sum_{k=1}^{p} |IPCA_{ik} \times EP_k| / \sum_{k=1}^{p} EP_k$$

where $WAAS_i$ is the weighted average of absolute scores of the *i*th genotype; PCA_{ik} is the score of the *i*th genotype in the *k*th IPCA; and EP_k is the explained variance of the *k*th IPCA for k = 1,2,...,p, considering p the number of significant PCAs.

Five simultaneous selection indexes (ssi) are also computed by summation of the ranks of the ASV, SIPC, EV and Za indexes and the ranks of the mean yields (Farshadfar, 2008), which results in ssiASV, ssiSIPC, ssiEV, ssiZa, and ssiWAAS, respectively.

Usage

AMMI_indexes(.data, order.y = NULL, level = 0.95)

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Arguments

.data An object of class waas or performs_ammi order.y A vector of the same length of x used to order the response variable. Each

element of the vector must be one of the 'h' or 'l'. If 'h' is used, the response variable will be ordered from maximum to minimum. If '1' is used then the response variable will be ordered from minimum to maximum. Use a comma-

separated vector of names. For example, order. y = c("h, h, 1, h, 1").

level The confidence level. Defaults to 0.95.

Value

A list where each element contains the result AMMI-based stability indexes for one variable.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Purchase, J.L., H. Hatting, and C.S. van Deventer. 2000. Genotype vs environment interaction of winter wheat (Triticum aestivum L.) in South Africa: II. Stability analysis of yield performance. South African J. Plant Soil 17:101-107. doi: 10.1080/02571862.2000.10634878

Sneller, C.H., L. Kilgore-Norquest, and D. Dombek. 1997. Repeatability of Yield Stability Statistics in Soybean. Crop Sci. 37:383-390. doi: 10.2135/cropsci1997.0011183X003700020013x

Zali, H., E. Farshadfar, S.H. Sabaghpour, and R. Karimizadeh. 2012. Evaluation of genotype vs environment interaction in chickpea using measures of stability from AMMI model. Ann. Biol. Res. 3:3126-3136.

Olivoto, T., A.D.C. L\'ucio, J.A.G. da silva, V.S. Marchioro, V.Q. de Souza, and E. Jost. 2019a. Mean performance and stability in multi-environment trials I: Combining features of AMMI and BLUP techniques. Agron. J. 111:2949-2960. doi: 10.2134/agronj2019.03.0220

```
library(metan)
model <- waas(data_ge,</pre>
               env = ENV.
               gen = GEN,
               rep = REP,
               resp = c(GY, HM),
               verbose = FALSE)
model_indexes <- AMMI_indexes(model)</pre>
# Alternatively (and more intuitively) using %>%
res_ind <- data_ge %>%
           waas(ENV, GEN, REP, c(GY, HM)) %>%
           AMMI_indexes()
```

10 Annicchiarico

Annicchiarico Annicchiarico's genotypic confidence index
--

Description

[Stable]

Stability analysis using the known genotypic confidence index (Annicchiarico, 1992).

Usage

```
Annicchiarico(.data, env, gen, rep, resp, prob = 0.25, verbose = TRUE)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s)
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
rep	The name of the column that contains the levels of the replications/blocks
resp	The response variable(s). To analyze multiple variables in a single procedure use, for example, resp = $c(var1, var2, var3)$.
prob	The probability of error assumed.
verbose	Logical argument. If verbose = FALSE the code will run silently.

Value

A list where each element is the result for one variable and contains the following data frames:

- **environments** Contains the mean, environmental index and classification as favorable and unfavorable environments.
- general Contains the genotypic confidence index considering all environments.
- favorable Contains the genotypic confidence index considering favorable environments.
- unfavorable Contains the genotypic confidence index considering unfavorable environments.

Author(s)

Tiago Olivoto, <tiagoolivoto@gmail.com>

References

Annicchiarico, P. 1992. Cultivar adaptation and recommendation from alfalfa trials in Northern Italy. J. Genet. Breed. 46:269-278.

See Also

```
superiority(), ecovalence(), ge_stats()
```

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Examples

anova_ind

Within-environment analysis of variance

Description

[Stable]

Performs a within-environment analysis of variance in randomized complete block or alpha-lattice designs and returns values such as Mean Squares, p-values, coefficient of variation, heritability, and accuracy of selection.

Usage

```
anova_ind(.data, env, gen, rep, resp, block = NULL, verbose = TRUE)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments. The analysis of variance is computed for each level of this factor.
gen	The name of the column that contains the levels of the genotypes.
rep	The name of the column that contains the levels of the replications/blocks.
resp	The response variable(s). To analyze multiple variables in a single procedure a vector of variables may be used. For example resp = c(var1, var2, var3).
block	Defaults to NULL. In this case, a randomized complete block design is considered. If block is informed, then a resolvable alpha-lattice design (Patterson and Williams, 1976) is employed. All effects, except the error, are assumed to be fixed.
verbose	Logical argument. If verbose = FALSE the code will run silently.

Value

A list where each element is the result for one variable containing (1) **individual**: A tidy tbl_df with the results of the individual analysis of variance with the following column names, and (2) **MSRatio**: The ratio between the higher and lower residual mean square. The following columns are returned, depending on the experimental design

• For analysis in alpha-lattice designs:

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- MEAN: The grand mean.
- DFG, DFCR, and DFIB_R, and DFE: The degree of freedom for genotype, complete replicates, incomplete blocks within replicates, and error, respectively.
- MSG, MSCR, MSIB_R: The mean squares for genotype, replicates, incomplete blocks within replicates, and error, respectively.
- FCG, FCR, FCIB_R: The F-calculated for genotype, replicates and incomplete blocks within replicates, respectively.
- PFG, PFCR, PFIB_R: The P-values for genotype, replicates and incomplete blocks within replicates, respectively.
- CV: coefficient of variation.
- h2: broad-sense heritability.
- AS: accuracy of selection (square root of h2)

• For analysis in randomized complete block design:

- MEAN: The grand mean.
- DFG, DFB, and DFE: The degree of freedom for genotype blocks, and error, respectively.
- MSG, MSB, and MSE: The mean squares for genotype blocks, and error, respectively.
- FCG and FCB: The F-calculated for genotype and blocks, respectively.
- PFG and PFB: The P-values for genotype and blocks, respectively.
- CV: coefficient of variation.
- h2: broad-sense heritability.
- AS: accuracy of selection (square root of h2)

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Patterson, H.D., and E.R. Williams. 1976. A new class of resolvable incomplete block designs. Biometrika 63:83-92.

anova_joint 13

anova_joint	Joint analysis of variance	

Description

[Stable]

Performs a joint analysis of variance to check for the presence of genotype-vs-environment interactions using both randomized complete block and alpha-lattice designs.

Usage

```
anova_joint(.data, env, gen, rep, resp, block = NULL, verbose = TRUE)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments. The analysis of variance is computed for each level of this factor.
gen	The name of the column that contains the levels of the genotypes.
rep	The name of the column that contains the levels of the replications/blocks.
resp	The response variable(s). To analyze multiple variables in a single procedure a vector of variables may be used. For example resp = c(var1, var2, var3).
block	Defaults to NULL. In this case, a randomized complete block design is considered. If block is informed, then a resolvable alpha-lattice design (Patterson and Williams, 1976) is employed. All effects, except the error, are assumed to be fixed.
verbose	Logical argument. If verbose = FALSE the code will run silently.

Value

A list where each element is the result for one variable containing the following objects:

- anova: The two-way ANOVA table
- model: The model of class 1m.
- augment: Information about each observation in the dataset. This includes predicted values in the fitted column, residuals in the resid column, standardized residuals in the stdres column, the diagonal of the 'hat' matrix in the hat, and standard errors for the fitted values in the se.fit column.
- **details:** A tibble with the following data: Ngen, the number of genotypes; OVmean, the grand mean; Min, the minimum observed (returning the genotype and replication/block); Max the maximum observed, MinGEN the loser winner genotype, MaxGEN, the winner genotype.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

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References

Patterson, H.D., and E.R. Williams. 1976. A new class of resolvable incomplete block designs. Biometrika 63:83-92.

See Also

```
get_model_data() anova_ind()
```

Examples

arrange_ggplot

Arrange separate ggplots into the same graphic

Description

[Experimental]

This is a wraper function around patchwork::wrap_plots() and patchwork::plot_annotation() to arrange ggplot2 objects.

Usage

```
arrange_ggplot(
  . . . ,
 nrow = NULL,
 ncol = NULL,
 widths = NULL,
 heights = NULL,
 guides = NULL,
 design = NULL,
 legend.position = "bottom",
  title = NULL,
 subtitle = NULL,
  caption = NULL,
  tag_levels = NULL,
  tag_prefix = NULL,
  tag_suffix = NULL,
  tag_sep = NULL,
```

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```
theme = NULL
)
```

Arguments

... multiple ggplots or a list containing ggplot objects.

nrow, ncol The number of rows and columns, respectively.

widths, heights

The relative widths and heights of each column and row in the grid. Will get

repeated to match the dimensions of the grid.

guides A string specifying how guides should be treated in the layout. Defaults to

'auto'. Other possible values are 'keep' and 'collect'. In this case, will

collect guides below to the given nesting level, removing duplicates.

design Specification of the location of areas in the layout.

legend.position

The position of the legends in the plot if guides = "collect" Default to 'bottom'.

title, subtitle, caption

Text strings to use for the various plot annotations.

tag_levels A character vector defining the enumeration format to use at each level. Possible

values are 'a' for lowercase letters, 'A' for uppercase letters, '1' for numbers, 'i' for lowercase Roman numerals, and 'I' for uppercase Roman numerals. It can also be a list containing character vectors defining arbitrary tag sequences. If any element in the list is a scalar and one of 'a', 'A', '1', 'i, or 'I', this level

will be expanded to the expected sequence.

tag_prefix, tag_suffix

Strings that should appear before or after the tag.

tag_sep A separator between different tag levels.

theme A ggplot theme specification to use for the plot. Only elements related to the

titles as well as plot margin and background is used.

Value

A patchwork object

as.lpcor

as.lpcor

Coerce to an object of class lpcor

Description

[Stable]

Functions to check if an object is of class 1pcor, or coerce it if possible.

Usage

```
as.lpcor(...)
```

Arguments

... A comma-separated list of matrices to be coerced to a list.

Value

An object of class 1pcor.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

barplots

Fast way to create bar plots

Description

[Stable]

- plot_bars() Creates a bar plot based on one categorical variable and one numeric variable. It can be used to show the results of a one-way trial with **qualitative treatments**.
- plot_factbars() Creates a bar plot based on two categorical variables and one numeric variable. It can be used to show the results of a two-way trial with **qualitative-qualitative treatment structure**.

Usage

```
plot_bars(
  .data,
  Х,
  у,
  order = NULL,
  y.lim = NULL,
  y.breaks = waiver(),
  y.expand = 0.05,
  y.contract = 0,
  xlab = NULL,
  ylab = NULL,
  n.dodge = 1,
  check.overlap = FALSE,
  color.bar = "black",
  fill.bar = "gray",
  lab.bar = NULL,
  lab.bar.hjust = 0.5,
  lab.bar.vjust = -0.5,
  lab.bar.angle = 0,
  size.text.bar = 5,
  values = FALSE,
  values.hjust = 0.5,
  values.vjust = 1.5,
  values.angle = 0,
  values.digits = 2,
  values.size = 4,
  lab.x.hjust = 0.5,
  lab.x.vjust = 1,
  lab.x.angle = 0,
  errorbar = TRUE,
  stat.erbar = "se",
  width.erbar = NULL,
  level = 0.95,
  invert = FALSE,
  width.bar = 0.9,
  size.line = 0.5,
```

```
size.text = 12,
  fontfam = "sans",
  na.rm = TRUE,
  verbose = FALSE,
  plot_theme = theme_metan()
plot_factbars(
  .data,
  ...,
  resp,
  y.lim = NULL,
  y.breaks = waiver(),
  y.expand = 0.05,
  y.contract = 0,
  xlab = NULL,
  ylab = NULL,
  n.dodge = 1,
  check.overlap = FALSE,
  lab.bar = NULL,
  lab.bar.hjust = 0.5,
  lab.bar.vjust = -0.5,
  lab.bar.angle = 0,
  size.text.bar = 5,
  values = FALSE,
  values.hjust = 0.5,
  values.vjust = 1.5,
  values.angle = 0,
  values.digits = 2,
  values.size = 4,
  lab.x.hjust = 0.5,
  lab.x.vjust = 1,
  lab.x.angle = 0,
  errorbar = TRUE,
  stat.erbar = "se",
  width.erbar = NULL,
  level = 0.95,
  invert = FALSE,
  col = TRUE,
  palette = "Spectral",
  width.bar = 0.9,
  legend.position = "bottom",
  size.line = 0.5,
  size.text = 12,
  fontfam = "sans",
  na.rm = TRUE,
  verbose = FALSE,
  plot_theme = theme_metan()
```

Arguments

. data The data set.

Argument valid for plot_bars() The variables to be mapped to the x and y х, у axes, respectively. Argument valid for plot_bars(). Controls the order of the factor in the x axis. order Defaults to the order of the factors in .data. Use order = "asce" or order = "desc" to reorder the labels to ascending or descending order, respectively, based on the values of the variable y. The range of y axis. Defaults to NULL (maximum and minimum values of the y.lim data set). New values can be inserted as y.lim = c(y.min, y.max). y.breaks The breaks to be plotted in the y-axis. Defaults to waiver(). authomatic breaks. The same arguments than x.breaks can be used. y.expand, y.contract A multiplication range expansion/contraction factor. y. expand expands the upper limit of the y escale, while y contract contracts the lower limit of the y scale. By default y.expand = 0.05 and y.contract = 0 produces a plot without spacing in the lower y limit and an expansion in the upper y limit. xlab, ylab The labels of the axes x and y, respectively. Defaults to NULL. n.dodge The number of rows that should be used to render the x labels. This is useful for displaying labels that would otherwise overlap. Silently remove overlapping labels, (recursively) prioritizing the first, last, and check.overlap middle labels. color.bar, fill.bar Argument valid for plot_bars(). The color and fill values of the bars. lab.bar A vector of characters to show in each bar. Defaults to NULL. lab.bar.hjust, lab.bar.vjust The horizontal and vertical adjust for the labels in the bar. Defaults to 0.5 and -0.5, respectively. lab.bar.angle The angle for the labels in the plot. Defaults to 0. Use in combination with lab.bar.hjust and lab.bar.vjust to best fit the labels in the plot. The size of the text in the bar labels. size.text.bar values Logical argument. Shows the values in the plot bar? Defaults to FALSE values.hjust, values.vjust The horizontal and vertical adjust for the values in the bar. Defaults to 0.5 and 1.5, respectively. If values = TRUE the values are shown bellow the error bar. values.angle The angle for the labels in the plot. Defaults to 0. Use in combination with values.hjust and values.vjust to best fit the values in the plot bar. values.digits The significant digits to show if values = TRUE. Defaults to 2. values.size The size of the text for values shown in the bars. Defaults to 3. lab.x.hjust, lab.x.vjust The horizontal and vertical adjust for the labels in the bar. Defaults to 0.5 and 1, respectively. The angle for the labels in x axis. Defaults to 0. Use in combination with lab.x.angle lab.x.hjust and lab.x.vjust to best fit the labels in the axis. errorbar Logical argument, set to TRUE. In this case, an error bar is shown. stat.erbar The statistic to be shown in the errorbar. Must be one of the stat.erbar = "se" (standard error, default), stat.erbar = "sd" (standard deviation), or stat.erbar = "ci" (confidence interval), based on the confidence level in the

argument level.

width.erbar	The width of the error bar. Defaults to 25% of width.bar.
level	The confidence level
invert	$Logical \ argument. \ If \ TRUE, \ rotate \ the \ plot \ in \ plot_bars() \ and \ invert \ the \ order \ of \ the \ factors \ in \ plot_factbars().$
width.bar	The width of the bars in the graph. Defaults to 0.9. Possible values are in the range 0 -1.
size.line	The size of the line in the bars. Default to 0.5.
size.text	The size of the text. Default to 12.
fontfam	The family of the font text. Defaults to "sans".
na.rm	Should 'NA' values be removed to compute the statistics? Defaults to true
verbose	Logical argument. If TRUE a tibble containing the mean, N, standard deviation, standard error of mean and confidence interval is returned.
plot_theme	The graphical theme of the plot. Default is plot_theme = theme_metan(). For more details, see ggplot2::theme().
•••	Argument valid for $plot_factbars()$. A comma-separated list of unquoted variable names. Sets the two variables to be mapped to the x axis.
resp	Argument valid for $plot_factbars()$. The response variable to be mapped to the y axis.
col	Logical argument valid for plot_factbars(). If FALSE, a gray scale is used.
palette	Argument valid for $plot_factbars()$ The color palette to be used. For more details, see ?scale_colour_brewer
legend.position	
	The position of the legend in the plot.

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

See Also

```
plot_lines(), plot_factlines()
```

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```
y.contract = -0.75, # contract the lower limit
errorbar = FALSE, # remove errorbar
color.bar = "red", # color of bars
fill.bar = alpha_color("cyan", 75), # create a transparent color
lab.bar = letters[1:13]) # add labels
arrange_ggplot(p1, p2)
```

bind_cv

Bind cross-validation objects

Description

[Stable]

Helper function that combines objects of class cv_ammi, cv_ammif or cv_blup. It is useful when looking for a boxplot containing the RMSPD values of those cross-validation procedures.

Usage

```
bind_cv(..., bind = "boot", sort = TRUE)
```

Arguments

... Input objects of class cv_ammi, cv_ammif or cv_blup.

bind What data should be used? To plot the RMSPD, use 'boot' (default). Use bind = 'means' to return the RMSPD mean for each model.

sort Used to sort the RMSPD mean in ascending order.

Value

An object of class cv_ammif. The results will depend on the argument bind. If bind = 'boot' then the RMSPD of all models in ... will be bind to a unique data frame. If bind = 'means' then the RMSPD mean of all models in ... will be bind to an unique data frame.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

22 blup_indexes

blup_indexes

Stability indexes based on a mixed-effect model

Description

[Stable]

- hmgv() Computes the harmonic mean of genotypic values (HMGV).
- rpgv() Computes the relative performance of the genotypic values (RPGV).
- hmrpgv() Computes the harmonic mean of the relative performance of genotypic values (HM-RPGV).
- blup_indexes() Is a wrapper around the above functions that also computes the WAASB index (Olivoto et al. 2019) if an object computed with waasb() is used as input data.
- Resende_indexes() [Deprecated] Use blup_indexes() instead.

Usage

```
hmgv(model)
rpgv(model)
hmrpgv(model)
Resende_indexes(model)
blup_indexes(model)
```

Arguments

model

An object of class waasb computed with waasb() or gamem_met().

Details

The indexes computed with this function have been used to select genotypes with stability performance in a mixed-effect model framework. Some examples are in Alves et al (2018), Azevedo Peixoto et al. (2018), Dias et al. (2018) and Colombari Filho et al. (2013).

The HMGV index is computed as

$$HMGV_i = \frac{E}{\sum_{j=1}^{E} \frac{1}{Gv_{ij}}}$$

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where E is the number of environments included in the analysis, Gv_{ij} is the genotypic value (BLUP) for the ith genotype in the jth environment.

The RPGV index is computed as

$$RPGV_i = \frac{1}{E} \sum_{j=1}^{E} Gv_{ij} / \mu_j$$

The HMRPGV index is computed as

$$HMRPGV_i = \frac{E}{\sum_{j=1}^{E} \frac{1}{Gv_{ij}/\mu_j}}$$

Value

A tibble containing the indexes.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Alves, R.S., L. de Azevedo Peixoto, P.E. Teodoro, L.A. Silva, E.V. Rodrigues, M.D.V. de Resende, B.G. Laviola, and L.L. Bhering. 2018. Selection of Jatropha curcas families based on temporal stability and adaptability of genetic values. Ind. Crops Prod. 119:290-293. doi: 10.1016/J.INDCROP.2018.04.029

Azevedo Peixoto, L. de, P.E. Teodoro, L.A. Silva, E.V. Rodrigues, B.G. Laviola, and L.L. Bhering. 2018. Jatropha half-sib family selection with high adaptability and genotypic stability. PLoS One 13:e0199880. doi: 10.1371/journal.pone.0199880

Colombari Filho, J.M., M.D.V. de Resende, O.P. de Morais, A.P. de Castro, E.P. Guimaraes, J.A. Pereira, M.M. Utumi, and F. Breseghello. 2013. Upland rice breeding in Brazil: a simultaneous genotypic evaluation of stability, adaptability and grain yield. Euphytica 192:117-129. doi: 10.1007/s1068101309222

Dias, P.C., A. Xavier, M.D.V. de Resende, M.H.P. Barbosa, F.A. Biernaski, R.A. Estopa. 2018. Genetic evaluation of Pinus taeda clones from somatic embryogenesis and their genotype x environment interaction. Crop Breed. Appl. Biotechnol. 18:55-64. doi: 10.1590/198470332018v18n1a8

Olivoto, T., A.D.C. L\'ucio, J.A.G. da silva, V.S. Marchioro, V.Q. de Souza, and E. Jost. 2019. Mean performance and stability in multi-environment trials I: Combining features of AMMI and BLUP techniques. Agron. J. 111:2949-2960. doi: 10.2134/agronj2019.03.0220

Resende MDV (2007) Matematica e estatistica na analise de experimentos e no melhoramento genetico. Embrapa Florestas, Colombo

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```
model_indexes <- blup_indexes(res_ind)
gmd(model_indexes)</pre>
```

can_corr

Canonical correlation analysis

Description

[Stable]

Performs canonical correlation analysis with collinearity diagnostic, estimation of canonical loads, canonical scores, and hypothesis testing for correlation pairs.

Usage

```
can_corr(
   .data,
   FG,
   SG,
   by = NULL,
   use = "cor",
   test = "Bartlett",
   prob = 0.05,
   center = TRUE,
   stdscores = FALSE,
   verbose = TRUE,
   collinearity = TRUE
)
```

Arguments

.data	The data to be analyzed. It can be a data frame (possible with grouped data passed from dplyr::group_by().
FG, SG	A comma-separated list of unquoted variable names that will compose the first (smallest) and second (highest) group of the correlation analysis, respectively. Select helpers are also allowed.
by	One variable (factor) to compute the function by. It is a shortcut to dplyr::group_by(). To compute the statistics by more than one grouping variable use that function.
use	The matrix to be used. Must be one of 'cor' for analysis using the correlation matrix (default) or 'cov' for analysis using the covariance matrix.
test	The test of significance of the relationship between the FG and SG. Must be one of the 'Bartlett' (default) or 'Rao'.
prob	The probability of error assumed. Set to 0.05.
center	Should the data be centered to compute the scores?
stdscores	Rescale scores to produce scores of unit variance?
verbose	Logical argument. If TRUE (default) then the results are shown in the console.
collinearity	Logical argument. If TRUE (default) then a collinearity diagnostic is performed for each group of variables according to Olivoto et al.(2017).

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Value

If .data is a grouped data passed from dplyr::group_by() then the results will be returned into a list-column of data frames.

- Matrix The correlation (or covariance) matrix of the variables
- MFG, MSG The correlation (or covariance) matrix for the variables of the first group or second group, respectively.
- MFG_SG The correlation (or covariance) matrix for the variables of the first group with the second group.
- Coef_FG, Coef_SG Matrix of the canonical coefficients of the first group or second group, respectively.
- Loads_FG, Loads_SG Matrix of the canonical loadings of the first group or second group, respectively.
- Score_FG, Score_SG Canonical scores for the variables in FG and SG, respectively.
- Crossload_FG, Crossload_FG Canonical cross-loadings for FG variables on the SG scores, and cross-loadings for SG variables on the FG scores, respectively.
- SigTest A dataframe with the correlation of the canonical pairs and hypothesis testing results.
- **collinearity** A list with the collinearity diagnostic for each group of variables.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Olivoto, T., V.Q. Souza, M. Nardino, I.R. Carvalho, M. Ferrari, A.J. Pelegrin, V.J. Szareski, and D. Schmidt. 2017. Multicollinearity in path analysis: a simple method to reduce its effects. Agron. J. 109:131-142. doi: 10.2134/agronj2016.04.0196

26 clustering

-			
clu	ste	ring	•

Clustering analysis

Description

[Stable]

Performs clustering analysis with selection of variables.

Usage

```
clustering(
   .data,
   ...,
   by = NULL,
   scale = FALSE,
   selvar = FALSE,
   verbose = TRUE,
   distmethod = "euclidean",
   clustmethod = "average",
   nclust = NA
)
```

Arguments

.data	The data to be analyzed. It can be a data frame, possible with grouped data passed from dplyr::group_by().
•••	The variables in .data to compute the distances. Set to NULL, i.e., all the numeric variables in .data are used.
by	One variable (factor) to compute the function by. It is a shortcut to dplyr::group_by(). To compute the statistics by more than one grouping variable use that function.
scale	Should the data be scaled before computing the distances? Set to FALSE. If TRUE, then, each observation will be divided by the standard deviation of the variable $Z_{ij} = X_{ij}/sd_j$
selvar	Logical argument, set to FALSE. If TRUE, then an algorithm for selecting variables is implemented. See the section Details for additional information.
verbose	Logical argument. If TRUE (default) then the results for variable selection are shown in the console.
distmethod	The distance measure to be used. This must be one of 'euclidean', 'maximum', 'manhattan', 'canberra', 'binary', 'minkowski', 'pearson', 'spearman', or 'kendall'. The last three are correlation-based distance.
clustmethod	The agglomeration method to be used. This should be one of 'ward.D', 'ward.D2', 'single', 'complete', 'average' (= UPGMA), 'mcquitty' (= WPGMA), 'median' (= WPGMC) or 'centroid' (= UPGMC).
nclust	The number of clusters to be formed. Set to NA

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Details

When selvar = TRUE a variable selection algorithm is executed. The objective is to select a group of variables that most contribute to explain the variability of the original data. The selection of the variables is based on eigenvalue/eigenvectors solution based on the following steps.

- 1. compute the distance matrix and the cophenetic correlation with the original variables (all numeric variables in dataset);
- 2. compute the eigenvalues and eigenvectors of the correlation matrix between the variables;
- 3. Delete the variable with the largest weight (highest eigenvector in the lowest eigenvalue);
- 4. Compute the distance matrix and cophenetic correlation with the remaining variables;
- Compute the Mantel's correlation between the obtained distances matrix and the original distance matrix;
- 6. Iterate steps 2 to 5 p 2 times, where p is the number of original variables.

At the end of the p - 2 iterations, a summary of the models is returned. The distance is calculated with the variables that generated the model with the largest cophenetic correlation. I suggest a careful evaluation aiming at choosing a parsimonious model, i.e., the one with the fewer number of variables, that presents acceptable cophenetic correlation and high similarity with the original distances.

Value

- data The data that was used to compute the distances.
- **cutpoint** The cutpoint of the dendrogram according to Mojena (1977).
- distance The matrix with the distances.
- de The distances in an object of class dist.
- hc The hierarchical clustering.
- Sqt The total sum of squares.
- tab A table with the clusters and similarity.
- clusters The sum of square and the mean of the clusters for each variable.
- cofgrap If selectvar = TRUE, then, cofpgrap is a ggplot2-based graphic showing the cophenetic correlation for each model (with different number of variables). Else, will be a NULL object.
- **statistics** If selectvar = TRUE, then, statistics shows the summary of the models fitted with different number of variables, including cophenetic correlation, Mantel's correlation with the original distances (all variables) and the p-value associated with the Mantel's test. Else, will be a NULL object.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Mojena, R. 2015. Hierarchical grouping methods and stopping rules: an evaluation. Comput. J. 20:359-363. doi: 10.1093/comjnl/20.4.359

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Examples

```
library(metan)
# All rows and all numeric variables from data
d1 <- clustering(data_ge2)</pre>
# Based on the mean for each genotype
mean_gen <-
 data_ge2 %>%
means_by(GEN) %>%
 column_to_rownames("GEN")
d2 <- clustering(mean_gen)</pre>
# Select variables for compute the distances
d3 <- clustering(mean_gen, selvar = TRUE)</pre>
# Compute the distances with standardized data
# Define 4 clusters
d4 <- clustering(data_ge,</pre>
                 by = ENV,
                  scale = TRUE,
                  nclust = 4)
```

coincidence_index

Computes the coincidence index of genotype selection

Description

[Stable]

Computes the coincidence index (Hamblin and Zimmermann, 1986) as follows:

$$CI = \frac{A - C}{M - C} \times 100$$

where A is the number of selected genotypes common to different methods; C is the number of expected genotypes selected by chance; and M is the number of genotypes selected according to the selection intensity.

Usage

```
coincidence_index(..., total, sel1 = NULL, sel2 = NULL)
```

Arguments

A comma-separated list of objects of class mgidi, mtsi fai_blup, or sh. When a model is informed, then the selected genotypes are extracted automatically.

The total number of genotypes in the study.

The selected genotypes by the method 1 and 2, respectively. Defaults to NULL.

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Value

A list with the following elements:

• **coincidence**: A data frame with the coincidence index, number of common genotypes and the list of common genotypes for each model combination.

- **coincidence_mat**: A matrix-like containing the coincidence index.
- **genotypes**: The number of common genotypes for all models, i.e., the insersection of the selected genotypes of all models

References

Hamblin, J., and M.J. de O. Zimmermann. 1986. Breeding Common Bean for Yield in Mixtures. p. 245-272. In Plant Breeding Reviews. John Wiley & Sons, Inc., Hoboken, NJ, USA.doi: 10.1002/9781118061015.ch8

Examples

```
sel1 <- paste("G", 1:30, sep = "")
sel2 <- paste("G", 16:45, sep = "")
coincidence_index(sel1 = sel1, sel2 = sel2, total = 150)</pre>
```

colindiag

Collinearity Diagnostics

Description

[Stable]

Perform a (multi)collinearity diagnostic of a correlation matrix of predictor variables using several indicators, as shown by Olivoto et al. (2017).

Usage

```
colindiag(.data, ..., by = NULL, n = NULL)
```

Arguments

.data	The data to be analyzed. It must be a symmetric correlation matrix, or a data frame, possible with grouped data passed from dplyr::group_by().
•••	Variables to use in the correlation. If is null then all the numeric variables from .data are used. It must be a single variable name or a comma-separated list of unquoted variables names.
by	One variable (factor) to compute the function by. It is a shortcut to dplyr::group_by(). To compute the statistics by more than one grouping variable use that function.
n	If a correlation matrix is provided, then n is the number of objects used to compute the correlation coefficients.

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Value

If .data is a grouped data passed from dplyr::group_by() then the results will be returned into a list-column of data frames.

- cormat A symmetric Pearson's coefficient correlation matrix between the variables
- corlist A hypothesis testing for each of the correlation coefficients
- evalevet The eigenvalues with associated eigenvectors of the correlation matrix
- VIF The Variance Inflation Factors, being the diagonal elements of the inverse of the correlation matrix.
- CN The Condition Number of the correlation matrix, given by the ratio between the largest and smallest eigenvalue.
- det The determinant of the correlation matrix.
- ncorhigh Number of correlation greather than 10.81.
- largest_corr The largest correlation (in absolute value) observed.
- smallest_corr The smallest correlation (in absolute value) observed.
- weight_var The variables with largest eigenvector (largest weight) in the eigenvalue of smallest value, sorted in decreasing order.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Olivoto, T., V.Q. Souza, M. Nardino, I.R. Carvalho, M. Ferrari, A.J. Pelegrin, V.J. Szareski, and D. Schmidt. 2017. Multicollinearity in path analysis: a simple method to reduce its effects. Agron. J. 109:131-142. doi: 10.2134/agronj2016.04.0196

comb_vars 31

comb	vars
COIIID	vais

Pairwise combinations of variables

Description

[Stable]

Pairwise combinations of variables that will be the result of a function applied to each combination.

Usage

```
comb_vars(.data, order = "first", FUN = "+", verbose = TRUE)
```

Arguments

.data	A matrix of data with, say, p columns.
order	The order on how the results will appear in the output. Default is order = 'first'. In this case, assuming that .data has four columns, namely, V1, V2, V3, V4, the order of columns in the output will be V1.V2, V1.V3, V1.V4, V2.V3, V2.V4, V3.V4. If order = 'second', the result will be then V1.V2, V1.V3, V2.V3, V1.V4, V2.V4, V3.V4.
FUN	The function that will be applied to each combination. The default is $+$, i.e., $V1 + V2$.
verbose	Logical argument. If verbose = FALSE the code will run silently.

Value

A data frame containing all possible combination of variables. Each combination is the result of the function in FUN applied to the two variables.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

32 correlated_vars

correlated_vars

Generate correlated variables

Description

[Experimental]

Generate correlated variables using a vector of know values and desired maximum and minimum correlations

Usage

```
correlated_vars(
   y,
   min_cor = -1,
   max_cor = 1,
   nvars,
   constant = NULL,
   operation = "*",
   x = NULL
)
```

Arguments

у	A vector to generate variables correlated with.
min_cor	The minimum desired correlation.
max_cor	The maximum desired correlation.
nvars	The number of variables.
constant	A constant. Use operation to define which operation is used.
operation	The operation to be applied to the constant value.
X	An optional vector of the same length of y. If not informed (default) then a normally distributed variable (mean $= 0$, $sd = 1$) will be used.

Value

A data frame with the y variable and the correlated variables.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

```
library(metan)
y <- rnorm(n = 10)
cor_vars <- correlated_vars(y, nvar = 6)
plot(cor_vars)</pre>
```

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corr_ci	corr	ci			
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Confidence interval for correlation coefficient

Description

[Stable]

Computes the half-width confidence interval for correlation coefficient using the nonparametric method proposed by Olivoto et al. (2018).

The half-width confidence interval is computed according to the following equation:

$$CI_w = 0.45304^r \times 2.25152 \times n^{-0.50089}$$

where n is the sample size and r is the correlation coefficient.

Usage

```
corr_ci(
   .data = NA,
   ...,
   r = NULL,
   n = NULL,
   by = NULL,
   sel.var = NULL,
   verbose = TRUE
)
```

Arguments

.data	The data to be analyzed. It can be a data frame (possible with grouped data passed from dplyr::group_by()) or a symmetric correlation matrix.
• • •	Variables to compute the confidence interval. If not informed, all the numeric variables from .data are used.
r	If data is not available, provide the value for correlation coefficient.
n	The sample size if data is a correlation matrix or if r is informed.
by	One variable (factor) to compute the function by. It is a shortcut to dplyr::group_by(). To compute the statistics by more than one grouping variable use that function.
sel.var	A variable to shows the correlation with. This will omit all the pairwise correlations that doesn't contain sel.var.
verbose	If verbose = TRUE then some results are shown in the console.

Value

A tibble containing the values of the correlation, confidence interval, upper and lower limits for all combination of variables.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

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References

Olivoto, T., A.D.C. Lucio, V.Q. Souza, M. Nardino, M.I. Diel, B.G. Sari, D.. K. Krysczun, D. Meira, and C. Meier. 2018. Confidence interval width for Pearson's correlation coefficient: a Gaussian-independent estimator based on sample size and strength of association. Agron. J. 110:1-8. doi: 10.2134/agronj2016.04.0196

Examples

corr_coef

Computes Pearson's correlation matrix with p-values

Description

Computes Pearson's correlation matrix with p-values

Usage

```
corr_coef(data, ..., verbose = TRUE)
```

Arguments

data The data set.

... Variables to use in the correlation. If no variable is informed all the numeric

variables from data are used.

verbose Logical argument. If verbose = FALSE the code is run silently.

Value

A list with the correlation coefficients and p-values

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

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Examples

```
library(metan)

# All numeric variables
all <- corr_coef(data_ge2)

# Select variables
sel <- corr_coef(data_ge2, EP, EL, CD, CL)
print(sel)</pre>
```

corr_plot

Visualization of a correlation matrix

Description

[Stable]

Graphical and numerical visualization of a correlation matrix

Usage

```
corr_plot(
  .data,
  . . . ,
  col.by = NULL,
  upper = "corr",
  lower = "scatter",
  decimal.mark = ".",
  axis.labels = FALSE,
  show.labels.in = "show",
  size.axis.label = 12,
  diag = TRUE,
  diag.type = "histogram",
  bins = 20,
  col.diag = "gray",
  alpha.diag = 1,
  col.up.panel = "gray",
  col.lw.panel = "gray",
  col.dia.panel = "gray",
  prob = 0.05,
  col.sign = "green",
  alpha.sign = 0.15,
  lab.position = "tr",
  progress = NULL,
  smooth = FALSE,
  col.smooth = "red",
  confint = TRUE,
  size.point = 1,
  shape.point = 19,
  alpha.point = 0.7,
```

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```
fill.point = NULL,
  col.point = "black",
  size.line = 0.5,
  minsize = 2,
  maxsize = 3,
  pan.spacing = 0.15,
  digits = 2,
  export = FALSE,
  file.type = "pdf",
  file.name = NULL,
  width = 8,
  height = 7,
  resolution = 300
)
```

Arguments

.data The data. Should, preferentially, contain numeric variables only. If .data has

factor-columns, these columns will be deleted with a warning message.

... Variables to use in the correlation. If no variable is informed all the numeric

variables from .data are used.

col.by A categorical variable to map the color of the points by. Defaults to NULL.

upper The visualization method for the upper triangular correlation matrix. Must be

one of 'corr' (numeric values), 'scatter' (the scatterplot for each pairwise

combination), or NULL to set a blank diagonal.

lower The visualization method for the lower triangular correlation matrix. Must be

one of 'corr' (numeric values), 'scatter' (the scatterplot for each pairwise

combination), or NULL to set a blank diagonal.

decimal.mark The decimal mark. Defaults to ".".

axis.labels Should the axis labels be shown in the plot? Set to FALSE.

show.labels.in Where to show the axis labels. Defaults to "show" bottom and left. Use "diag"

to show the labels on the diagonal. In this case, the diagonal layer (boxplot,

density or histogram) will be overwritten.

size.axis.label

The size of the text for axis labels if axis.labels = TRUE. Defaults to 12.

diag Should the diagonal be shown?

diag.type The type of plot to show in the diagonal if diag TRUE. It must be one of the

'histogram' (to show an histogram), 'density' to show the Kernel density, or

'boxplot' (to show a boxplot).

bins The number of bins, Defaults to 20.

col.diag If diag = TRUE then diagcol is the color for the distribution. Set to gray.

alpha.diag Alpha-transparency scale (0-1) to make the diagonal plot transparent. 0 = fully

transparent; 1 = full color. Set to 0.15

col.up.panel, col.lw.panel, col.dia.panel

The color for the upper, lower, and diagonal panels, respectively. Set to 'gray'.

prob The probability of error. Significant correlations will be highlighted with '',

", and " (0.05, 0.01, and 0.001, respectively). Scatterplots with significant

correlations may be color-highlighted.

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col.sign	The color that will highlight the significant correlations. Set to 'green'.
alpha.sign	Alpha-transparency scale (0-1) to make the plot area transparent. $0 = \text{fully transparent}$; $1 = \text{full color}$. Set to 0.15
lab.position	The position that the labels will appear. Set to 'tr', i.e., the legends will appear in the top and right of the plot. Other allowed options are 'tl' (top and left), 'br' (bottom and right), 'bl' (bottom and left).
progress	NULL (default) for a progress bar in interactive sessions with more than 15 plots, TRUE for a progress bar, FALSE for no progress bar.
smooth	Should a linear smooth line be shown in the scatterplots? Set to FALSE.
col.smooth	The color for the smooth line.
confint	Should a confidence band be shown with the smooth line? Set to TRUE.
size.point	The size of the points in the plot. Set to 0.5 .
shape.point	The shape of the point, set to 1.
alpha.point	Alpha-transparency scale (0-1) to make the points transparent. $0 = \text{fully transparent}$; $1 = \text{full color}$. Set to 0.7
fill.point	The color to fill the points. Valid argument if points are between 21 and 25.
col.point	The color for the edge of the point, set to black.
size.line	The size of the line (smooth and diagonal).
minsize	The size of the letter that will represent the smallest correlation coefficient.
maxsize	The size of the letter that will represent the largest correlation coefficient.
pan.spacing	The space between the panels. Set to 0.15.
digits	The number of digits to show in the plot.
export	Logical argument. If TRUE, then the plot is exported to the current directory.
file.type	The format of the file if export = TRUE. Set to 'pdf'. Other possible values are *.tiff using file.type = 'tiff'.
file.name	The name of the plot when exported. Set to NULL, i.e., automatically.
width	The width of the plot, set to 8.
height	The height of the plot, set to 7.
resolution	The resolution of the plot if file.type = 'tiff' is used. Set to 300 (300 dpi).

Value

An object of class gg, ggmatrix.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

```
library(metan)
dataset <- data_ge2 %>% select_cols(1:7)
# Default plot setting
corr_plot(dataset)
# Chosing variables to be correlated
```

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```
corr_plot(dataset, PH, EH, EL)
# Axis labels, similar to the function pairs()
# Gray scale
corr_plot(dataset, PH, EH, EL,
          shape.point = 19,
          size.point = 2,
          alpha.point = 0.5,
          alpha.diag = 0,
          pan.spacing = 0,
          col.sign = 'gray',
          alpha.sign = 0.3,
          axis.labels = TRUE)
corr_plot(dataset, PH, EH, EL,
          prob = 0.01,
          shape.point = 21,
          col.point = 'black',
          fill.point = 'orange',
          size.point = 2,
          alpha.point = 0.6,
          maxsize = 4,
          minsize = 2,
          smooth = TRUE,
          size.line = 1,
          col.smooth = 'black',
          col.sign = 'cyan',
          col.up.panel = 'black',
          col.lw.panel = 'black',
          col.dia.panel = 'black',
          pan.spacing = 0,
          lab.position = 'tl')
```

corr_ss

Sample size planning for a desired Pearson's correlation confidence interval

Description

[Stable]

Find the required (sufficient) sample size for computing a Pearson correlation coefficient with a desired confidence interval (Olivoto et al., 2018) as follows

$$n = \left[\frac{CI_w}{0.45304^r \times 2.25152}\right]^{-0.50089}$$

where CI_w is desired confidence interval and r is the correlation coefficient.

Usage

```
corr_ss(r, CI, verbose = TRUE)
```

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Arguments

r The magnitude of the correlation coefficient.

CI The half-width for confidence interval at p < 0.05.

verbose Logical argument. If verbose = FALSE the code is run silently.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Olivoto, T., A.D.C. Lucio, V.Q. Souza, M. Nardino, M.I. Diel, B.G. Sari, D.. K. Krysczun, D. Meira, and C. Meier. 2018. Confidence interval width for Pearson's correlation coefficient: a Gaussian-independent estimator based on sample size and strength of association. Agron. J. 110:1-8. doi: 10.2134/agronj2016.04.0196

Examples

```
corr_ss(r = 0.60, CI = 0.1)
```

corr_stab_ind

Correlation between stability indexes

Description

[Stable]

Computes the Spearman's rank correlation between the parametric and nonparametric stability indexes computed with the function ge_stats().

Usage

```
corr_stab_ind(x, stats = "all", plot = TRUE, ...)
```

Arguments

X	An object of class ge_stats.
stats	The statistics to compute the correlation. See the section Details for more information.
plot	Plot the heat map with the correlations? Defaults to TRUE.
	Other arguments to be passed to the function plot.corr_coef().

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Details

The argument stats is used to chose the statistics to show the ranks. Allowed values are "all" (All statistics, default), "par" (Parametric statistics), "nonpar" (Non-parametric statistics), "ammi" (AMMI-based stability statistics), or the following values that can be combined into comma-separated character vector. "Y" (Response variable), "Var" (Genotype's variance), "Shukla" (Shukla's variance), "Wi_g", "Wi_t", "Wi_u" (Annichiarrico's genotypic confidence index for all, favorable and unfavorable environments, respectively), "Ecoval" (Wricke's ecovalence), "Sij" (Deviations from the joint-regression analysis), "R2" (R-squared from the joint-regression analysis), "ASV" (AMMIstability value), "SIPC" (sum of the absolute values of the IPCA scores), "EV" (Average of the squared eigenvector values), "ZA" (Absolute values of the relative contributions of the IPCAs to the interaction), "WAAS" (Weighted Average of Absolute Scores), "HMGV" (Harmonic mean of the genotypic value), "RPGV" (Relative performance of the genotypic values), "HMRPGV" (Harmonic mean of the relative performance of the genotypic values), "Pi_a", "Pi_f", "Pi_u" (Superiority indexes for all, favorable and unfavorable environments, respectively), "Gai" (Geometric adaptability index), "\$1" (mean of the absolute rank differences of a genotype over the n environments), "\$2" (variance among the ranks over the k environments), "S3" (sum of the absolute deviations), "S6" (relative sum of squares of rank for each genotype), "N1", "N2", "N3", "N4" (Thennarasu"s statistics)).

Value

A list with the data (ranks) correlation, p-values and a heat map showing the correlation coefficients.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
model <- ge_stats(data_ge, ENV, GEN, REP, GY)
a <- corr_stab_ind(model)
b <- corr_stab_ind(model, stats = "ammi")
c <- corr_stab_ind(model, stats = c("ASV, Sij, R2, WAAS, N1"))</pre>
```

covcor_design

Variance-covariance matrices for designed experiments

Description

[Stable]

Compute variance-covariance and correlation matrices using data from a designed (RCBD or CRD) experiment.

Usage

```
covcor_design(.data, gen, rep, resp, design = "RCBD", by = NULL, type = NULL)
```

covcor_design 41

Arguments

.data	The data to be analyzed. It can be a data frame, possible with grouped data passed from dplyr::group_by().
gen	The name of the column that contains the levels of the genotypes.
rep	The name of the column that contains the levels of the replications/blocks.
resp	The response variables. For example $resp = c(var1, var2, var3)$.
design	The experimental design. Must be RCBD or CRD.
by	One variable (factor) to compute the function by. It is a shortcut to dplyr::group_by(). To compute the statistics by more than one grouping variable use that function.
type	What the matrices should return? Set to NULL, i.e., a list of matrices is returned. The argument type allow the following values 'pcor', 'gcor', 'rcor', (which will return the phenotypic, genotypic and residual correlation matrices, respectively) or 'pcov', 'gcov', 'rcov' (which will return the phenotypic, genotypic and residual variance-covariance matrices, respectively). Alternatively, it is possible to get a matrix with the means of each genotype in each trait, by using type = 'means'.

Value

An object of class covcor_design containing the following items:

- **geno_cov** The genotypic covariance.
- phen_cov The phenotypic covariance.
- resi_cov The residual covariance.
- **geno_cor** The phenotypic correlation.
- phen_cor The phenotypic correlation.
- resi_cor The residual correlation.

If .data is a grouped data passed from dplyr::group_by() then the results will be returned into a list-column of data frames.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

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cv_ammi

Cross-validation procedure

Description

[Stable]

Cross-validation for estimation of AMMI models

THe original dataset is split into two datasets: training set and validation set. The 'training' set has all combinations (genotype x environment) with N-1 replications. The 'validation' set has the remaining replication. The splitting of the dataset into modeling and validation sets depends on the design informed. For Completely Randomized Block Design (default), and alpha-lattice design (declaring block arguments), complete replicates are selected within environments. The remained replicate serves as validation data. If design = 'RCD' is informed, completely randomly samples are made for each genotype-by-environment combination (Olivoto et al. 2019). The estimated values considering naxis-Interaction Principal Component Axis are compared with the 'validation' data. The Root Mean Square Prediction Difference (RMSPD) is computed. At the end of boots, a list is returned.

IMPORTANT: If the data set is unbalanced (i.e., any genotype missing in any environment) the function will return an error. An error is also observed if any combination of genotype-environment has a different number of replications than observed in the trial.

Usage

```
cv_ammi(
   .data,
   env,
   gen,
   rep,
   resp,
   block = NULL,
   naxis = 2,
   nboot = 200,
   design = "RCBD",
   verbose = TRUE
)
```

Arguments

. data The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).

env The name of the column that contains the levels of the environments.

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cv ammı	43
cv_amm	7.5

gen	The name of the column that contains the levels of the genotypes.
rep	The name of the column that contains the levels of the replications/blocks. AT LEAST THREE REPLICATES ARE REQUIRED TO PERFORM THE CROSS-VALIDATION.
resp	The response variable.
block	Defaults to NULL. In this case, a randomized complete block design is considered. If block is informed, then a resolvable alpha-lattice design (Patterson and Williams, 1976) is employed. All effects, except the error, are assumed to be fixed.
naxis	The number of axis to be considered for estimation of GE effects.
nboot	The number of resamples to be used in the cross-validation. Defaults to 200.
design	The experimental design. Defaults to RCBD (Randomized complete Block Design). For Completely Randomized Designs inform design = $'CRD'$.
verbose	A logical argument to define if a progress bar is shown. Default is TRUE.

Value

An object of class cv_ammi with the following items: * **RMSPD**: A vector with nboot-estimates of the Root Mean Squared Prediction Difference between predicted and validating data.

- RMSPDmean: The mean of RMSPDmean estimates.
- **Estimated**: A data frame that contain the values (predicted, observed, validation) of the last loop.
- Modeling: The dataset used as modeling data in the last loop
- Testing: The dataset used as testing data in the last loop.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Olivoto, T., A.D.C. L\'ucio, J.A.G. da silva, V.S. Marchioro, V.Q. de Souza, and E. Jost. 2019. Mean performance and stability in multi-environment trials I: Combining features of AMMI and BLUP techniques. Agron. J. 111:2949-2960. doi: 10.2134/agronj2019.03.0220

Patterson, H.D., and E.R. Williams. 1976. A new class of resolvable incomplete block designs. Biometrika 63:83-92.

See Also

```
cv_ammif(), cv_blup()
```

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```
nboot = 5,
naxis = 2)
```

cv_ammif

Cross-validation procedure

Description

[Stable]

Cross-validation for estimation of all AMMI-family models

cv_ammif provides a complete cross-validation of replicate-based data using AMMI-family models. By default, the first validation is carried out considering the AMMIF (all possible axis used). Considering this model, the original dataset is split up into two datasets: training set and validation set. The 'training' set has all combinations (genotype x environment) with N-1 replications. The 'validation' set has the remaining replication. The splitting of the dataset into modeling and validation sets depends on the design informed. For Completely Randomized Block Design (default), and alpha-lattice design (declaring block arguments), complete replicates are selected within environments. The remained replicate serves as validation data. If design = 'RCD' is informed, completely randomly samples are made for each genotype-by-environment combination (Olivoto et al. 2019). The estimated values for each member of the AMMI-family model are compared with the 'validation' data. The Root Mean Square Prediction Difference (RMSPD) is computed. At the end of boots, a list is returned.

IMPORTANT: If the data set is unbalanced (i.e., any genotype missing in any environment) the function will return an error. An error is also observed if any combination of genotype-environment has a different number of replications than observed in the trial.

Usage

```
cv_ammif(
   .data,
   env,
   gen,
   rep,
   resp,
   nboot = 200,
   block,
   design = "RCBD",
   verbose = TRUE
)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.

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rep	The name of the column that contains the levels of the replications/blocks. AT LEAST THREE REPLICATES ARE REQUIRED TO PERFORM THE CROSS-VALIDATION.
resp	The response variable.
nboot	The number of resamples to be used in the cross-validation. Defaults to 200.
block	Defaults to NULL. In this case, a randomized complete block design is considered. If block is informed, then a resolvable alpha-lattice design (Patterson and Williams, 1976) is employed. All effects, except the error, are assumed to be fixed.
design	The experimental design used in each environment. Defaults to RCBD (Randomized complete Block Design). For Completely Randomized Designs inform design = 'CRD'.
verbose	A logical argument to define if a progress bar is shown. Default is TRUE.

Value

An object of class cv_ammif with the following items:

- RMSPD: A vector with nboot-estimates of the Root Mean Squared Prediction Difference between predicted and validating data.
- RMSPDmean: The mean of RMSPDmean estimates.
- **Estimated**: A data frame that contain the values (predicted, observed, validation) of the last loop.
- Modeling: The dataset used as modeling data in the last loop
- **Testing**: The dataset used as testing data in the last loop.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Patterson, H.D., and E.R. Williams. 1976. A new class of resolvable incomplete block designs. Biometrika 63:83-92.

See Also

```
cv_ammi(), cv_blup()
```

 cv_blup

cv_blup	Cross-validation procedure	

Description

[Stable]

Cross-validation for blup prediction.

This function provides a cross-validation procedure for mixed models using replicate-based data. By default, complete blocks are randomly selected within each environment. In each iteration, the original dataset is split up into two datasets: training and validation data. The 'training' set has all combinations (genotype x environment) with R - 1 replications. The 'validation' set has the remaining replication. The estimated values are compared with the 'validation' data and the Root Means Square Prediction Difference (Olivoto et al. 2019) is computed. At the end of boots, a list is returned.

Usage

```
cv_blup(
   .data,
   env,
   gen,
   rep,
   resp,
   block = NULL,
   nboot = 200,
   random = "gen",
   verbose = TRUE
)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
rep	The name of the column that contains the levels of the replications/blocks. AT LEAST THREE REPLICATES ARE REQUIRED TO PERFORM THE CROSS-VALIDATION.
resp	The response variable.
block	Defaults to NULL. In this case, a randomized complete block design is considered. If block is informed, then a resolvable alpha-lattice design (Patterson and Williams, 1976) is employed. See how fixed and random effects are considered, see the section Details .
nboot	The number of resamples to be used in the cross-validation. Defaults to 200
random	The effects of the model assumed to be random. See Details for more information.
verbose	A logical argument to define if a progress bar is shown. Default is TRUE.

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Details

Six models may be fitted depending upon the values in block and random arguments.

• Model 1: block = NULL and random = "gen" (The default option). This model considers a Randomized Complete Block Design in each environment assuming genotype and genotype-environment interaction as random effects. Environments and blocks nested within environments are assumed to fixed factors.

- Model 2: block = NULL and random = "env". This model considers a Randomized Complete Block Design in each environment treating environment, genotype-environment interaction, and blocks nested within environments as random factors. Genotypes are assumed to be fixed factors.
- Model 3: block = NULL and random = "all". This model considers a Randomized Complete Block Design in each environment assuming a random-effect model, i.e., all effects (genotypes, environments, genotype-vs-environment interaction and blocks nested within environments) are assumed to be random factors.
- Model 4: block is not NULL and random = "gen". This model considers an alpha-lattice design in each environment assuming genotype, genotype-environment interaction, and incomplete blocks nested within complete replicates as random to make use of inter-block information (Mohring et al., 2015). Complete replicates nested within environments and environments are assumed to be fixed factors.
- Model 5: block is not NULL and random = "env". This model considers an alpha-lattice design in each environment assuming genotype as fixed. All other sources of variation (environment, genotype-environment interaction, complete replicates nested within environments, and incomplete blocks nested within replicates) are assumed to be random factors.
- **Model 6:** block is not NULL and random = "all". This model considers an alpha-lattice design in each environment assuming all effects, except the intercept, as random factors.

IMPORTANT: An error is returned if any combination of genotype-environment has a different number of replications than observed in the trial.

Value

An object of class cv_blup with the following items: * **RMSPD**: A vector with nboot-estimates of the root mean squared prediction difference between predicted and validating data. * **RMSPDmean** The mean of RMSPDmean estimates.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Olivoto, T., A.D.C. L\'ucio, J.A.G. da silva, V.S. Marchioro, V.Q. de Souza, and E. Jost. 2019. Mean performance and stability in multi-environment trials I: Combining features of AMMI and BLUP techniques. Agron. J. 111:2949-2960. doi: 10.2134/agronj2019.03.0220

Patterson, H.D., and E.R. Williams. 1976. A new class of resolvable incomplete block designs. Biometrika 63:83-92.

Mohring, J., E. Williams, and H.-P. Piepho. 2015. Inter-block information: to recover or not to recover it? TAG. Theor. Appl. Genet. 128:1541-54. doi: 10.1007/s0012201525300

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See Also

```
cv_ammi(), cv_ammif()
```

Examples

data_alpha

Data from an alpha lattice design

Description

Alpha lattice design of spring oats

Format

A tibble with 72 observations on the following 5 variables.

- PLOT Plot number
- REP Replicate code
- BLOCK Incomplete block code
- **GEN** Genotype code
- YIELD Observed dry matter yield (tonnes/ha)

Details

A spring oats trial grown in Craibstone. There were 24 varieties in 3 replicates, each consisting of 6 incomplete blocks of 4 plots. Planted in a resolvable alpha design. The plots were laid out in a single line.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Source

J. A. John & E. R. Williams (1995). Cyclic and computer generated designs, Chapman and Hall, London. Page 146.

data_g 49

data_g

Single maize trial

Description

This dataset contain data on 15 traits assessed in 13 maize hybrids. The experimental design was a RCBD with 3 blocks and 1 replications per block. It is used as an example in the function gamem() of the **metan** package.

Format

A tibble with 39 observations on the following 17 variables.

- GEN A factor with 13 levels; each level represents one maize hybrid.
- **REP** A factor with 3 levels; each level represents one replication/block.
- PH Plant height, in cm.
- EH Ear height, in cm.
- **EP** Ear position, i.e., the ratio EH/PH.
- EL Ear length, in cm.
- ED Ear diameter, in mm.
- CL Cob length, in cm.
- CD Cob diameter, in mm.
- CW Cob weight, in g.
- KW Kernel weight, in cm.
- NR Number of rows.
- NKR Number of kernels per row.
- CDED Cob diameter / Ear diameter ratio.
- **PERK** Percentage of kernels.
- · TKW Thousand-kernel weight
- NKE Number of kernels per row.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Source

Personal data

50 data_ge2

data_ge

Multi-environment trial of oat

Description

This dataset contain data on two variables assessed in 10 genotypes growing in in 11 environments. The experimental design was a RCBD with 3 replicates(blocks). This data provide examples for several functions of **metan** package.

Format

A tibble with 420 observations on the following 5 variables.

- ENV A factor with 14 levels; each level represents one cultivation environment.
- GEN A factor with 10 levels; each level represents one genotype.
- **REP** A factor with 3 levels; each level represents one replication/block.
- GY A continuous variable (grain yield) observed in each plot.
- HM A continuous variable (hectoliter mass) observed in each plot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Source

Personal data

data_ge2

Multi-environment trial of maize

Description

This dataset contain data on 15 traits assessed in 13 maize hybrids growing in 4 environments. The experimental design was a RCBD with 3 blocks and 1 replications per block. It may be used as example in several functions of **metan** package.

Format

A tibble with 156 observations on the following 18 variables.

- ENV A factor with 4 levels; each level represents one cultivation environment.
- GEN A factor with 13 levels; each level represents one maize hybrid.
- **REP** A factor with 3 levels; each level represents one replication/block.
- PH Plant height, in cm.
- EH Ear height, in cm.
- **EP** Ear position, i.e., the ratio EH/PH.
- EL Ear length, in cm.

data_simula 51

- ED Ear diameter, in mm.
- CL Cob length, in cm.
- CD Cob diameter, in mm.
- CW Cob weight, in g.
- KW Kernel weight, in cm.
- NR Number of rows.
- NKR Number of kernels per row.
- CDED Cob diameter / Ear diameter ratio.
- PERK Percentage of kernels.
- TKW Thousand-kernel weight
- NKE Number of kernels per row.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Source

Personal data

data_simula

Simulate genotype and genotype-environment data

Description

[Experimental]

- g_simula() simulate replicated genotype data.
- ge_simula() simulate replicated genotype-environment data.

Usage

```
ge_simula(
    ngen,
    nenv,
    nrep,
    nvars = 1,
    gen_eff = 20,
    env_eff = 15,
    rep_eff = 5,
    ge_eff = 10,
    res_eff = 5,
    intercept = 100,
    seed = NULL
)

g_simula(
    ngen,
```

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```
nrep,
nvars = 1,
gen_eff = 20,
rep_eff = 5,
res_eff = 5,
intercept = 100,
seed = NULL
)
```

Arguments

ngen	The number of genotypes.
nenv	The number of environments.
nrep	The number of replications.
nvars	The number of traits.
gen_eff	The genotype effect.
env_eff	The environment effect
rep_eff	The replication effect
ge_eff	The genotype-environment interaction effect.
res_eff	The residual effect. The effect is sampled from a normal distribution with zero mean and standard deviation equal to res_eff. Be sure to change res_eff when changin the intercept scale.
intercept	The intercept.
seed	The seed.

Details

The functions simulate genotype or genotype-environment data given a desired number of genotypes, environments and effects. All effects are sampled from an uniform distribution. For example, given 10 genotypes, and $gen_eff = 30$, the genotype effects will be sampled as runif(10,min = -30,max = 30). Use the argument seed to ensure reproducibility. If more than one trait is used (nvars > 1), the effects and seed can be passed as a numeric vector. Single numeric values will be recycled with a warning when more than one trait is used.

Value

A data frame with the simulated traits

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

desc_stat 53

```
inspect(gen_data, plot = TRUE)
aov(V1 ~ GEN + REP, data = gen_data) %>% anova()
# Genotype-environment data
# 5 genotypes, 3 environments, 4 replicates and 2 traits
df <-
ge_simula(ngen = 5,
          nenv = 3,
          nrep = 4,
          nvars = 2,
          seed = 1)
ge_plot(df, ENV, GEN, V1)
aov(V1 ~ ENV*GEN + ENV/REP, data = df) %>% anova()
# Change genotype effect (trait 1 with fewer differences among genotypes)
# Define different intercepts for the two traits
df2 <-
ge\_simula(ngen = 10,
          nenv = 3,
          nrep = 4,
          nvars = 2,
          gen_eff = c(1, 50),
          intercept = c(80, 1500),
          seed = 1)
ge_plot(df2, ENV, GEN, V2)
```

 $desc_stat$

Descriptive statistics

Description

[Stable]

- desc_stat() Computes the most used measures of central tendency, position, and dispersion.
- desc_wider() is useful to put the variables in columns and grouping variables in rows. The table is filled with a statistic chosen with the argument stat.

Usage

```
desc_stat(
   .data = NULL,
   ...,
   by = NULL,
   stats = "main",
   hist = FALSE,
   level = 0.95,
   digits = 4,
   na.rm = FALSE,
   verbose = TRUE,
   plot_theme = theme_metan()
)

desc_wider(.data, which)
```

54 desc_stat

Arguments

.data

The data to be analyzed. It can be a data frame (possible with grouped data passed from dplyr::group_by() or a numeric vector. For desc_wider() .data is an object of class desc_stat.

. . .

A single variable name or a comma-separated list of unquoted variables names. If no variable is informed, all the numeric variables from .data will be used. Select helpers are allowed.

by

One variable (factor) to compute the function by. It is a shortcut to dplyr::group_by(). To compute the statistics by more than one grouping variable use that function.

stats

The descriptive statistics to show. This is used to filter the output after computation. Defaults to "main" (cv, max, mean median, min, sd.amo, se, ci). Other allowed values are "all" to show all the statistics, "robust" to show robust statistics, "quantile" to show quantile statistics, or chose one (or more) of the following:

- "av.dev": average deviation.
- "ci": 95 percent confidence interval of the mean.
- "cv": coefficient of variation.
- "iqr": interquartile range.
- "gmean": geometric mean.
- "hmean": harmonic mean.
- "Kurt": kurtosis.
- "mad": median absolute deviation.
- "max": maximum value.
- "mean": arithmetic mean.
- "median": median.
- "min": minimum value.
- "n": the length of the data.
- "n.valid": The valid (Not NA) number of elements
- "n.missing": The number of missing values
- "n.unique": The length of unique elements.
- "ps": the pseudo-sigma (iqr / 1.35).
- "q2.5", "q25", "q75", "q97.5": the percentile 2.5\ quartile, third quartile, and percentile 97.5\
- range: The range of data).
- "sd.amo", "sd.pop": the sample and population standard deviation.
- "se": the standard error of the mean.
- "skew": skewness.
- "sum". the sum of the values.
- "sum.dev": the sum of the absolute deviations.
- "sum. sq. dev": the sum of the squared deviations.
- "n.valid": The size of sample with valid number (not NA).
- "var.amo", "var.pop": the sample and population variance.

Use a names to select the statistics. For example, stats = c("median,mean,cv,n"). Note that the statistic names **are not** case-sensitive. Both comma or space can be used as separator.

hist

Logical argument defaults to FALSE. If hist = TRUE then a histogram is created for each selected variable.

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level	The confidence level to compute the confidence interval of mean. Defaults to 0.95.
digits	The number of significant digits.
na.rm	Logical. Should missing values be removed? Defaults to FALSE.
verbose	Logical argument. If verbose = FALSE the code is run silently.
plot_theme	The graphical theme of the plot. Default is plot_theme = theme_metan(). For more details, see ggplot2::theme().
which	A statistic to fill the table.

Value

- desc_stats() returns a tibble with the statistics in the columns and variables (with possible grouping factors) in rows.
- desc_wider() returns a tibble with variables in columns and grouping factors in rows.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

```
library(metan)
# Example 1: main statistics (coefficient of variation, maximum, #
# mean, median, minimum, sample standard deviation, standard
# error and confidence interval of the mean) for all numeric
# variables in data
desc_stat(data_ge2)
#Example 2: robust statistics using a numeric vector as input #
# data
vect <- data_ge2$TKW</pre>
desc_stat(vect, stats = "robust")
# Example 3: Select specific statistics. In this example, NAs #
# are removed before analysis with a warning message
desc_stat(c(12, 13, 19, 21, 8, NA, 23, NA),
      stats = c('mean, se, cv, n, n.valid'),
      na.rm = TRUE)
# Example 4: Select specific variables and compute statistics by#
# levels of a factor variable (GEN)
stats <-
 desc_stat(data_ge2,
        EP, EL, EH, ED, PH, CD,
        by = GEN)
```

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```
stats
```

doo

Alternative to dplyr::do for doing anything

Description

Provides an alternative to the dplyr:do() using nest(), mutate() and map() to apply a function to a grouped data frame.

Usage

```
doo(.data, .fun, ..., unnest = TRUE)
```

Arguments

.data a (grouped) data frame
.fun A function, formula, or atomic vector.
... Additional arguments passed on to .fun
unnest Logical argument defaults to TRUE to control if results of .fun should be unnested. Valid only if the result is of class data.frame or tbl_df.

Details

If the applied function returns a data frame, then the output will be automatically unnested. Otherwise, the output includes the grouping variables and a column named "data", which is a "list-columns" containing the results for group combinations.

Value

a data frame

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

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Examples

```
library(metan)
# Head the first two lines of each environment
data_ge2 %>%
  group_by(ENV) %>%
  doo(~head(., 2))

# Genotype analysis for each environment using 'gafem()'
# variable PH
data_ge2 %>%
  group_by(ENV) %>%
  doo(~gafem(., GEN, REP, PH, verbose = FALSE))
```

ecovalence

Stability analysis based on Wricke's model

Description

[Stable]

The function computes the ecovalence (Wricke, 1965) for stability analysis.

Usage

```
ecovalence(.data, env, gen, rep, resp, verbose = TRUE)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
rep	The name of the column that contains the levels of the replications/blocks.
resp	The response variable(s). To analyze multiple variables in a single procedure use, for example, $resp = c(var1, var2, var3)$.
verbose	Logical argument. If verbose = FALSE the code will run silently.

Value

An object of class ecovalence containing the results for each variable used in the argument resp.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Wricke, G. 1965. Zur berechnung der okovalenz bei sommerweizen und hafer. Z. Pflanzenzuchtg 52:127-138.

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Examples

env_dissimilarity

Dissimilarity between environments

Description

[Stable]

Computes the dissimilarity between environments based on several approaches. See the section **details** for more details.

Usage

```
env_dissimilarity(.data, env, gen, rep, resp)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
rep	The name of the column that contains the levels of the replications/blocks.
resp	The response variable(s). To analyze multiple variables in a single procedure a vector of variables may be used. For example resp = c(var1, var2, var3). Select helpers are also allowed.

Details

Roberteson (1959) proposed the partition of the mean square of the genotype-environment interaction (MS_GE) into single (S) and complex (C) parts, where $S=\frac{1}{2}(\sqrt{Q1}-\sqrt{Q2})^2)$ and $C=(1-r)\sqrt{Q1-Q2}$, being r the correlation between the genotype's average in the two environments; and Q1 and Q2 the genotype mean square in the environments 1 and 2, respectively. Cruz and Castoldi (1991) proposed a new decomposition of the MS_GE, in which the complex part is given by $C=\sqrt{(1-r)^3\times Q1\times Q2}$.

Value

A list with the following matrices:

• SPART_CC: The percentage of the single (non cross-over) part of the interaction between genotypes and pairs of environments according to the method proposed by Cruz and Castoldi (1991).

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• CPART_CC: The percentage of the complex (cross-over) part of the interaction between genotypes and pairs of environments according to the method proposed by Cruz and Castoldi (1991).

- SPART_RO: The percentage of the single (non cross-over) part of the interaction between genotypes and pairs of environments according to the method proposed by Robertson (1959).
- CPART_RO: The percentage of the complex (cross-over) part of the interaction between genotypes and pairs of environments according to the method proposed by Robertson (1959).
- MSGE: Interaction mean square between genotypes and pairs of environments.
- SSGE: Interaction sum of square between genotypes and pairs of environments.
- correlation: Correlation coefficients between genotypes's average in each pair of environment.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Cruz, C.D., Castoldi, F. (1991). Decomposicao da interacao genotipos x ambientes em partes simples e complexa. Ceres, 38:422-430.

Robertson, A. (1959). Experimental design on the measurement of heritabilities and genetic correlations. biometrical genetics. New York: Pergamon Press.

Examples

```
library(metan)
mod <- env_dissimilarity(data_ge, ENV, GEN, REP, GY)
print(mod)</pre>
```

env_stratification

Environment stratification

Description

[Stable]

Computes environment stratification based on factor analysis.

Usage

```
env_stratification(
   .data,
   env,
   gen,
   resp,
   use = "complete.obs",
   mineval = 1,
   verbose = TRUE
)
```

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Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s)
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
resp	The response variable(s). To analyze multiple variables in a single procedure use, for example, $resp = c(var1, var2, var3)$.
use	The method for computing covariances in the presence of missing values. Defaults to complete.obs, i.e., missing values are handled by casewise deletion.
mineval	The minimum value so that an eigenvector is retained in the factor analysis.
verbose	Logical argument. If verbose = FALSE the code will run silently.

Value

An object of class env_stratification which is a list with one element per analyzed trait. For each trait, the following values are given.

- data The genotype-environment means.
- cormat: The correlation matrix among the environments.
- PCA: The eigenvalues and explained variance.
- FA: The factor analysis.
- env_strat: The environmental stratification.
- mega_env_code: The environments within each mega-environment.
- mega_env_stat: The statistics for each mega-environment.
- KMO: The result for the Kaiser-Meyer-Olkin test.
- MSA: The measure of sampling adequacy for individual variable.
- communalities_mean: The communalities' mean.
- initial_loadings: The initial loadings.

Author(s)

Tiago Olivoto, <tiagoolivoto@gmail.com>

References

Murakami, D.M.D., and C.D.C. Cruz. 2004. Proposal of methodologies for environment stratification and analysis of genotype adaptability. Crop Breed. Appl. Biotechnol. 4:7-11.

See Also

```
env_dissimilarity()
```

fai_blup 61

Examples

fai_blup

Multi-trait selection index

Description

[Stable]

Multitrait index based on factor analysis and ideotype-design proposed by Rocha et al. (2018).

Usage

```
fai_blup(
   .data,
   use_data = "blup",
   DI = NULL,
   UI = NULL,
   SI = 15,
   mineval = 1,
   verbose = TRUE
)
```

Arguments

verbose

.data	An object of class waasb or a two-way table with genotypes in the rows and traits in columns. In the last case the row names must contain the genotypes names.
use_data	Define which data to use If .data is an object of class gamem. Defaults to "blup" (the BLUPs for genotypes). Use "pheno" to use phenotypic means instead BLUPs for computing the index.
DI, UI	A vector of the same length of .data to construct the desirable (DI) and undesirable (UI) ideotypes. For each element of the vector, allowed values are 'max', 'min', 'mean', or a numeric value. Use a comma-separated vector of text. For example, DI = $c("max,max,min,min")$. By default, DI is set to "max" for all traits and UI is set to "min" for all traits.
SI	An integer (0-100). The selection intensity in percentage of the total number of genotypes. Defaults to 15.
mineval	The minimum value so that an eigenvector is retained in the factor analysis.

Logical value. If TRUE some results are shown in console.

find_outliers

Value

An object of class fai_blup with the following items:

- data The data (BLUPS) used to compute the index.
- eigen The eigenvalues and explained variance for each axis.
- FA The results of the factor analysis.
- canonical_loadings The canonical loadings for each factor retained.
- FAI A list with the FAI-BLUP index for each ideotype design.
- sel_dif_trait A list with the selection differential for each ideotype design.
- sel_gen The selected genotypes.
- ideotype_construction A list with the construction of the ideotypes.
- total_gain A list with the total gain for variables to be increased or decreased.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Rocha, J.R.A.S.C.R, J.C. Machado, and P.C.S. Carneiro. 2018. Multitrait index based on factor analysis and ideotype-design: proposal and application on elephant grass breeding for bioenergy. GCB Bioenergy 10:52-60. doi: 10.1111/gcbb.12443

Examples

find_outliers

Find possible outliers in a dataset

Description

[Stable]

Find possible outliers in the dataset.

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Usage

```
find_outliers(
   .data = NULL,
   var = NULL,
   by = NULL,
   plots = FALSE,
   coef = 1.5,
   verbose = TRUE,
   plot_theme = theme_metan()
)
```

Arguments

.data	The data to be analyzed. Must be a dataframe or an object of class split_factors.
var	The variable to be analyzed.
by	One variable (factor) to compute the function by. It is a shortcut to dplyr::group_by(). To compute the statistics by more than one grouping variable use that function.
plots	If TRUE, then histograms and boxplots are shown.
coef	The multiplication coefficient, defaults to 1.5. For more details see ?boxplot.stat.
verbose	If verbose = TRUE then some results are shown in the console.
plot_theme	The graphical theme of the plot. Default is plot_theme = theme_metan(). For more details, see ggplot2::theme().

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
find_outliers(data_ge2, var = PH, plots = TRUE)
# Find outliers within each environment
find_outliers(data_ge2, var = PH, by = ENV)
```

Fox

Fox's stability function

Description

[Stable]

Performs a stability analysis based on the criteria of Fox et al. (1990), using the statistical "TOP third" only. A stratified ranking of the genotypes at each environment is done. The proportion of locations at which the genotype occurred in the top third are expressed in the output.

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Usage

```
Fox(.data, env, gen, resp, verbose = TRUE)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
resp	The response variable(s). To analyze multiple variables in a single procedure use, for example, $resp = c(var1, var2, var3)$.
verbose	Logical argument. If verbose = FALSE the code will run silently.

Value

An object of class Fox, which is a list containing the results for each variable used in the argument resp. For each variable, a tibble with the following columns is returned.

- **GEN** the genotype's code.
- mean the mean for the response variable.
- TOP The proportion of locations at which the

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Fox, P.N., B. Skovmand, B.K. Thompson, H.J. Braun, and R. Cormier. 1990. Yield and adaptation of hexaploid spring triticale. Euphytica 47:57-64. doi: 10.1007/BF00040364.

Examples

```
library(metan)
out <- Fox(data_ge2, ENV, GEN, PH)
print(out)</pre>
```

gafem Genotype analysis by fixed-effect models	
--	--

Description

[Stable]

One-way analysis of variance of genotypes conducted in both randomized complete block and alpha-lattice designs.

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Usage

```
gafem(
   .data,
   gen,
   rep,
   resp,
   block = NULL,
   by = NULL,
   prob = 0.05,
   verbose = TRUE
)
```

Arguments

.data

	response variable(s).
gen	The name of the column that contains the levels of the genotypes, that will be treated as random effect.
rep	The name of the column that contains the levels of the replications (assumed to be fixed).
resp	The response variable(s). To analyze multiple variables in a single procedure a vector of variables may be used. For example resp = c(var1, var2, var3). Select helpers are also allowed.
block	Defaults to NULL. In this case, a randomized complete block design is considered. If block is informed, then a resolvable alpha-lattice design (Patterson and Williams, 1976) is employed. All effects, except the error, are assumed to be fixed. Use the function gamem() to analyze a one-way trial with mixed-effect models.
by	One variable (factor) to compute the function by. It is a shortcut to <pre>dplyr::group_by()</pre> . This is especially useful, for example, when the researcher want to fit a fixed-effect model for each environment. In this case, an object of class gafem_grouped is returned. <pre>mgidi()</pre> can then be used to compute the mgidi index within each environment.

The dataset containing the columns related to, Genotypes, replication/block and

Details

prob

verbose

gafem analyses data from a one-way genotype testing experiment. By default, a randomized complete block design is used according to the following model:

Logical argument. If verbose = FALSE the code are run silently.

The error probability. Defaults to 0.05.

$$Y_{ij} = m + g_i + r_j + e_{ij}$$

where Y_{ij} is the response variable of the ith genotype in the jth block; m is the grand mean (fixed); g_i is the effect of the jth replicate; and e_{ij} is the random error.

When block is informed, then a resolvable alpha design is implemented, according to the following model:

$$Y_{ijk} = m + g_i + r_j + b_{jk} + e_{ijk}$$

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where where y_{ijk} is the response variable of the *i*th genotype in the *k*th block of the *j*th replicate; m is the intercept, t_i is the effect for the *i*th genotype r_j is the effect of the *j*th replicate, b_{jk} is the effect of the *k*th incomplete block of the *j*th replicate, and e_{ijk} is the plot error effect corresponding to y_{ijk} . All effects, except the random error are assumed to be fixed.

Value

A list where each element is the result for one variable containing the following objects:

- anova: The one-way ANOVA table.
- model: The model with of lm.
- augment: Information about each observation in the dataset. This includes predicted values in the fitted column, residuals in the resid column, standardized residuals in the stdres column, the diagonal of the 'hat' matrix in the hat, and standard errors for the fitted values in the se.fit column.
- hsd: The Tukey's 'Honest Significant Difference' for genotype effect.
- **details:** A tibble with the following data: Ngen, the number of genotypes; OVmean, the grand mean; Min, the minimum observed (returning the genotype and replication/block); Max the maximum observed, MinGEN the loser winner genotype, MaxGEN, the winner genotype.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Patterson, H.D., and E.R. Williams. 1976. A new class of resolvable incomplete block designs. Biometrika 63:83-92.

See Also

```
get_model_data() gamem()
```

```
library(metan)
# RCBD
rcbd <- gafem(data_g,</pre>
             gen = GEN,
             rep = REP,
              resp = c(PH, ED, EL, CL, CW))
# Fitted values
get_model_data(rcbd)
# ALPHA-LATTICE DESIGN
alpha <- gafem(data_alpha,</pre>
               gen = GEN,
               rep = REP,
               block = BLOCK,
               resp = YIELD)
# Fitted values
get_model_data(alpha)
```

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gai

Geometric adaptability index

Description

[Stable]

Performs a stability analysis based on the geometric mean (GAI), according to the following model (Mohammadi and Amri, 2008):

$$GAI = \sqrt[E]{\bar{Y}_1 + \bar{Y}_2 + \dots + \bar{Y}_i}$$

where \bar{Y}_1 , \bar{Y}_2 , and \bar{Y}_i are the mean yields of the first, second and *i*-th genotypes across environments, and E is the number of environments

Usage

```
gai(.data, env, gen, rep = "deprecated", resp, verbose = TRUE)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
rep	[Deprecated]
resp	The response variable(s). To analyze multiple variables in a single procedure use, for example, $resp = c(var1, var2, var3)$.
verbose	Logical argument. If verbose = FALSE the code will run silently.

Value

An object of class gai, which is a list containing the results for each variable used in the argument resp. For each variable, a tibble with the following columns is returned.

- **GEN** the genotype's code.
- GAI Geometric adaptability index
- GAI_R The rank for the GAI value.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Mohammadi, R., & Amri, A. (2008). Comparison of parametric and non-parametric methods for selecting stable and adapted durum wheat genotypes in variable environments. Euphytica, 159(3), 419-432. doi: 10.1007/s1068100796006.

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Examples

```
library(metan)
out <- gai(data_ge2, ENV, GEN, REP, c(EH, PH, EL, CD, ED, NKE))</pre>
```

gamem

Genotype analysis by mixed-effect models

Description

[Stable]

Analysis of genotypes in single experiments using mixed-effect models with estimation of genetic parameters.

Usage

```
gamem(
   .data,
   gen,
   rep,
   resp,
   block = NULL,
   by = NULL,
   prob = 0.05,
   verbose = TRUE
)
```

Arguments

verbose

.data	The dataset containing the columns related to, Genotypes, replication/block and response variable(s).
gen	The name of the column that contains the levels of the genotypes, that will be treated as random effect.
rep	The name of the column that contains the levels of the replications (assumed to be fixed).
resp	The response variable(s). To analyze multiple variables in a single procedure a vector of variables may be used. For example resp = c(var1, var2, var3). Select helpers are also allowed.
block	Defaults to NULL. In this case, a randomized complete block design is considered. If block is informed, then an alpha-lattice design is employed considering block as random to make use of inter-block information, whereas the complete replicate effect is always taken as fixed, as no inter-replicate information was to be recovered (Mohring et al., 2015).
by	One variable (factor) to compute the function by. It is a shortcut to dplyr::group_by(). This is especially useful, for example, when the researcher want to fit a mixed-effect model for each environment. In this case, an object of class gamem_grouped is returned. mgidi() can then be used to compute the mgidi index within each environment.
prob	The probability for estimating confidence interval for BLUP's prediction.

Logical argument. If verbose = FALSE the code are run silently.

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Details

gamem analyses data from a one-way genotype testing experiment. By default, a randomized complete block design is used according to the following model:

$$Y_{ij} = m + g_i + r_j + e_{ij}$$

where Y_{ij} is the response variable of the ith genotype in the jth block; m is the grand mean (fixed); g_i is the effect of the ith genotype (assumed to be random); r_j is the effect of the jth replicate (assumed to be fixed); and e_{ij} is the random error.

When block is informed, then a resolvable alpha design is implemented, according to the following model:

$$Y_{ijk} = m + g_i + r_j + b_{jk} + e_{ijk}$$

where where y_{ijk} is the response variable of the *i*th genotype in the *k*th block of the *j*th replicate; m is the intercept, t_i is the effect for the *i*th genotype r_j is the effect of the *j*th replicate, b_{jk} is the effect of the *k*th incomplete block of the *j*th replicate, and e_{ijk} is the plot error effect corresponding to y_{ijk} .

Value

An object of class gamem or gamem_grouped, which is a list with the following items for each element (variable):

- fixed: Test for fixed effects.
- random: Variance components for random effects.
- LRT: The Likelihood Ratio Test for the random effects.
- BLUPgen: The estimated BLUPS for genotypes
- ranef: The random effects of the model
- modellme The mixed-effect model of class 1merMod.
- residuals The residuals of the mixed-effect model.
- model_lm The fixed-effect model of class 1m.
- residuals_lm The residuals of the fixed-effect model.
- **Details:** A tibble with the following data: Ngen, the number of genotypes; OVmean, the grand mean; Min, the minimum observed (returning the genotype and replication/block); Max the maximum observed, MinGEN the winner genotype, MaxGEN, the loser genotype.
- **ESTIMATES:** A tibble with the values:
 - Gen_var, the genotypic variance and ;
 - rep:block_var block-within-replicate variance (if an alpha-lattice design is used by informing the block in block);
 - Res_var, the residual variance;
 - Gen (%), rep:block (%), and Res (%) the respective contribution of variance components to the phenotypic variance;
 - H2, broad-sense heritability;
 - h2mg, heritability on the entry-mean basis;
 - Accuracy, the accuracy of selection (square root of h2mg);
 - CVg, genotypic coefficient of variation;
 - CVr, residual coefficient of variation;
 - CV ratio, the ratio between genotypic and residual coefficient of variation.
- formula The formula used to fit the mixed-model.

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Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Mohring, J., E. Williams, and H.-P. Piepho. 2015. Inter-block information: to recover or not to recover it? TAG. Theor. Appl. Genet. 128:1541-54. doi: 10.1007/s0012201525300

See Also

```
get_model_data() waasb()
```

```
library(metan)
# fitting the model considering an RCBD
# Genotype as random effects
rcbd <- gamem(data_g,</pre>
             gen = GEN,
             rep = REP,
             resp = c(PH, ED, EL, CL, CW, KW, NR, TKW, NKE))
# Likelihood ratio test for random effects
get_model_data(rcbd, "lrt")
# Variance components
get_model_data(rcbd, "vcomp")
# Genetic parameters
get_model_data(rcbd, "genpar")
# random effects
get_model_data(rcbd, "ranef")
# Predicted values
predict(rcbd)
# fitting the model considering an alpha-lattice design
# Genotype and block-within-replicate as random effects
# Note that block effect was now informed.
alpha <- gamem(data_alpha,</pre>
               gen = GEN,
               rep = REP,
               block = BLOCK,
               resp = YIELD)
# Genetic parameters
get_model_data(alpha, "genpar")
# Random effects
get_model_data(alpha, "ranef")
```

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gamem_met

Genotype-environment analysis by mixed-effect models

Description

[Stable]

Genotype analysis in multi-environment trials using mixed-effect or random-effect models.

The nature of the effects in the model is chosen with the argument random. By default, the experimental design considered in each environment is a randomized complete block design. If block is informed, a resolvable alpha-lattice design (Patterson and Williams, 1976) is implemented. The following six models can be fitted depending on the values of random and block arguments.

- Model 1: block = NULL and random = "gen" (The default option). This model considers a Randomized Complete Block Design in each environment assuming genotype and genotype-environment interaction as random effects. Environments and blocks nested within environments are assumed to fixed factors.
- Model 2: block = NULL and random = "env". This model considers a Randomized Complete Block Design in each environment treating environment, genotype-environment interaction, and blocks nested within environments as random factors. Genotypes are assumed to be fixed factors.
- Model 3: block = NULL and random = "all". This model considers a Randomized Complete Block Design in each environment assuming a random-effect model, i.e., all effects (genotypes, environments, genotype-vs-environment interaction and blocks nested within environments) are assumed to be random factors.
- Model 4: block is not NULL and random = "gen". This model considers an alpha-lattice design in each environment assuming genotype, genotype-environment interaction, and incomplete blocks nested within complete replicates as random to make use of inter-block information (Mohring et al., 2015). Complete replicates nested within environments and environments are assumed to be fixed factors.
- Model 5: block is not NULL and random = "env". This model considers an alpha-lattice design in each environment assuming genotype as fixed. All other sources of variation (environment, genotype-environment interaction, complete replicates nested within environments, and incomplete blocks nested within replicates) are assumed to be random factors.
- **Model 6:** block is not NULL and random = "all". This model considers an alpha-lattice design in each environment assuming all effects, except the intercept, as random factors.

Usage

```
gamem_met(
    .data,
    env,
    gen,
    rep,
    resp,
    block = NULL,
    by = NULL,
    random = "gen",
    prob = 0.05,
    verbose = TRUE
)
```

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Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
rep	The name of the column that contains the levels of the replications/blocks.
resp	The response variable(s). To analyze multiple variables in a single procedure a vector of variables may be used. For example resp = c(var1, var2, var3).
block	Defaults to NULL. In this case, a randomized complete block design is considered. If block is informed, then an alpha-lattice design is employed considering block as random to make use of inter-block information, whereas the complete replicate effect is always taken as fixed, as no inter-replicate information was to be recovered (Mohring et al., 2015).
by	One variable (factor) to compute the function by. It is a shortcut to dplyr::group_by(). This is especially useful, for example, when the researcher want to analyze environments within mega-environments. In this case, an object of class waasb_grouped is returned.
random	The effects of the model assumed to be random. Defaults to random = "gen". See Details to see the random effects assumed depending on the experimental design of the trials.
prob	The probability for estimating confidence interval for BLUP's prediction.
verbose	Logical argument. If verbose = FALSE the code will run silently.

Value

An object of class waasb with the following items for each variable:

- **fixed** Test for fixed effects.
- random Variance components for random effects.
- LRT The Likelihood Ratio Test for the random effects.
- **BLUPgen** The random effects and estimated BLUPS for genotypes (If random = "gen" or random = "all")
- **BLUPenv** The random effects and estimated BLUPS for environments, (If random = "env" or random = "all").
- **BLUPint** The random effects and estimated BLUPS of all genotypes in all environments.
- MeansGxE The phenotypic means of genotypes in the environments.
- modellme The mixed-effect model of class 1merMod.
- residuals The residuals of the mixed-effect model.
- model lm The fixed-effect model of class 1m.
- **residuals_lm** The residuals of the fixed-effect model.
- **Details** A list summarizing the results. The following information are shown: Nenv, the number of environments in the analysis; Ngen the number of genotypes in the analysis; Mean the grand mean; SE the standard error of the mean; SD the standard deviation. CV the coefficient of variation of the phenotypic means, estimating WAASB, Min the minimum value observed (returning the genotype and environment), Max the maximum value observed (returning the genotype and environment); MinENV the environment with the lower mean, MaxENV the environment with the larger mean observed, MinGEN the genotype with the lower mean, MaxGEN the genotype with the larger.

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• **ESTIMATES** A tibble with the genetic parameters (if random = "gen" or random = "all") with the following columns: Phenotypic variance the phenotypic variance; Heritability the broad-sense heritability; GEr2 the coefficient of determination of the interaction effects; h2mg the heritability on the mean basis; Accuracy the selective accuracy; rge the genotype-environment correlation; CVg the genotypic coefficient of variation; CVr the residual coefficient of variation; CV ratio the ratio between genotypic and residual coefficient of variation.

• formula The formula used to fit the mixed-model.

Author(s)

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References

Olivoto, T., A.D.C. L\'ucio, J.A.G. da silva, V.S. Marchioro, V.Q. de Souza, and E. Jost. 2019. Mean performance and stability in multi-environment trials I: Combining features of AMMI and BLUP techniques. Agron. J. 111:2949-2960. doi: 10.2134/agronj2019.03.0220

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Patterson, H.D., and E.R. Williams. 1976. A new class of resolvable incomplete block designs. Biometrika 63:83-92.

See Also

```
mtsi() waas() get_model_data() plot_scores()
```

```
library(metan)
# Example 1: Analyzing all numeric variables assuming genotypes #
# as random effects
model <- gamem_met(data_ge,</pre>
            env = ENV,
            gen = GEN,
            rep = REP,
           resp = everything())
# Distribution of random effects (first variable)
plot(model, type = "re")
# Genetic parameters
get_model_data(model, "genpar")
#
# Example 2: Unbalanced trials
                                          #
# assuming all factors as random effects
un_data <- data_ge %>%
        remove_rows(1:3) %>%
        droplevels()
model2 <- gamem_met(un_data,</pre>
```

74 get_corvars

```
env = ENV,
    gen = GEN,
    rep = REP,
    random = "all",
    resp = GY)
get_model_data(model2)
```

get_corvars

Generate normal, correlated variables

Description

[Stable]

Given the mean and desired correlations, generate normal, correlated variables.

Usage

```
get_corvars(n = 10, mu, sigma, tol = 1e-06, seed = NULL)
```

Arguments

n	The number of samples required.
mu	A vector with the means for the variables.
sigma	A symmetric, positive-definite matrix with the (co)variance or correlation matrix of the variables.
tol	Tolerance (relative to largest variance) for numerical lack of positive-definiteness in sigma.
seed	An integer value interpreted as seed.

Value

A tibble containing the simulated data.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

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get_covmat

Generate a covariance matrix

Description

[Stable]

Given the variances and desired correlations, generate a covariance matrix

Usage

```
get_covmat(cormat, var)
```

Arguments

cormat

A symmetric matrix with desired correlations.

var

A numeric vector with variances. It must have length equal to the number of

elements in the diagonal of cormat.

Value

A (co)variance matrix

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

get_dist

Get a distance matrix

Description

[Stable]

Get the distance matrices from objects fitted with the function clustering(). This is especially useful to get distance matrices from several objects to be further analyzed using pairs_mantel().

Usage

```
get_dist(..., digits = 2)
```

Arguments

Object(s) of class clustering.] . . .

The number of significant figures. Defaults to 2. digits

Value

A list of class clustering.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
d <- data_ge2 %>%
      means_by(GEN) %>%
      column_to_rownames("GEN") %>%
      clustering()
get_dist(d)
```

get_model_data

Get data from a model easily

Description

[Experimental]

Usage

```
get_model_data(x, what = NULL, type = "GEN", verbose = TRUE)
gmd(x, what = NULL, type = "GEN", verbose = TRUE)
sel_gen(x)
```

Arguments

X	An object created with the functions AMMI_indexes(), anova_ind(), anova_joint(), can_corr() ecovalence(), Fox(), gai(), gamem(), gafem(), ge_acv(), ge_means(), ge_reg(), gytb(), mgidi(), performs_ammi(), blup_indexes(), Shukla(), superiority(), waas() or waasb().
what	What should be captured from the model. See more in section Details .

Chose if the statistics must be show by genotype (type = "GEN", default) or type

environment (TYPE = "ENV"), when possible.

verbose Logical argument. If verbose = FALSE the code will run silently.

Details

• get_model_data() Easily get data from some objects generated in the **metan** package such as the WAASB and WAASBY indexes (Olivoto et al., 2019a, 2019b) BLUPs, variance components, details of AMMI models and AMMI-based stability statistics.

- gmd() Is a shortcut to get_model_data.
- sel_gen() Extracts the selected genotypes by a given index.

Bellow are listed the options allowed in the argument what depending on the class of the object

Objects of class AMMI_indexes:

- "ASV" AMMI stability value.
- "EV" Averages of the squared eigenvector values.
- "SIPC" Sums of the absolute value of the IPCA scores.
- "WAAS" Weighted average of absolute scores (default).
- "ZA" Absolute value of the relative contribution of IPCAs to the interaction.

Objects of class anova_ind:

- "MEAN" The mean value of the variable
- "DFG", "DFB", "DFCR", "DFIB_R", "DFE". The degree of freedom for genotypes, blocks (randomized complete block design), complete replicates, incomplete blocks within replicates (alpha-lattice design), and error, respectively.
- "MSG", "FCG", "PFG" The mean square, F-calculated and P-values for genotype effect, respectively.
- "MSB", "FCB", "PFB" The mean square, F-calculated and P-values for block effect in randomized complete block design.
- "MSCR", "FCR", "PFCR" The mean square, F-calculated and P-values for complete replicates in alpha lattice design.
- "MSIB_R", "FCIB_R", "PFIB_R" The mean square, F-calculated and P-values for incomplete blocks within complete replicates, respectively (for alpha lattice design only).
- "MSE" The mean square of error.
- "CV" The coefficient of variation.
- "h2" The broad-sence heritability
- "AS" The accurary of selection (square root of h2).
- "FMAX" The Hartley's test (the ratio of the largest MSE to the smallest MSE).

Objects of class anova_joint or gafem:

- "Y" The observed values.
- "h2" The broad-sense heritability.
- "Sum Sq" Sum of squares.
- "Mean Sq" Mean Squares.
- "F value" F-values.
- "Pr(>F)" P-values.
- ".fitted" Fitted values (default).
- ".resid" Residuals.
- ".stdresid" Standardized residuals.

- ".se.fit" Standard errors of the fitted values.
- "details" Details.

Objects of class Annicchiarico and Schmildt:

- "Sem_rp" The standard error of the relative mean performance (Schmildt).
- "Mean_rp" The relative performance of the mean.
- "rank" The rank for genotypic confidence index.
- "Wi" The genotypic confidence index.

Objects of class can_corr:

- "coefs" The canonical coefficients (default).
- "loads" The canonical loadings.
- "crossloads" The canonical cross-loadings.
- "canonical" The canonical correlations and hypothesis testing.

Objects of class ecovalence:

- "Ecoval" Ecovalence value (default).
- "Ecov_perc" Ecovalence in percentage value.
- "rank" Rank for ecovalence.

Objects of class fai_blup: See the **Value** section of fai_blup() to see valid options for what argument.

Objects of class ge_acv:

- "ACV" The adjusted coefficient of variation (default).
- "ACV_R" The rank for adjusted coefficient of variation.

Objects of class ge_polar:

- "POLAR" The Power Law Residuals (default).
- "POLAR_R" The rank for Power Law Residuals.

Objects of class ge_reg:

- GEN: the genotypes.
- b0 and b1 (default): the intercept and slope of the regression, respectively.
- t(b1=1): the calculated t-value
- pval_t: the p-value for the t test.
- s2di the deviations from the regression (stability parameter).
- F(s2di=0): the F-test for the deviations.
- pval_f: the p-value for the F test;
- RMSE the root-mean-square error.
- R2 the determination coefficient of the regression.

Objects of class ge_effects:

• For objects of class ge_effects no argument what is required.

Objects of class ge_means:

- "ge_means" Genotype-environment interaction means (default).
- "env_means" Environment means.
- "gen_means" Genotype means.

Objects of class gge:

- "scores" The scores for genotypes and environments for all the analyzed traits (default).
- "exp_var" The eigenvalues and explained variance.
- "projection" The projection of each genotype in the AEC coordinates in the stability GGE plot

Objects of class gytb:

- "gyt" Genotype by yield*trait table (Default).
- "stand_gyt" The standardized (zero mean and unit variance) Genotype by yield*trait table.
- "si" The superiority index (sum standardized value across all yield*trait combinations).

Objects of class mgidi: See the **Value** section of mgidi() to see valid options for what argument. **Objects of class** mtsi: See the **Value** section of mtsi() to see valid options for what argument.

Objects of class Shukla:

- "rMean" Rank for the mean.
- "ShuklaVar" Shukla's stablity variance (default).
- "rShukaVar" Rank for Shukla's stablity variance.
- "ssiShukaVar" Simultaneous selection index.

Objects of class sh: See the **Value** section of Smith_Hazel() to see valid options for what argument.

Objects of class Fox:

• "TOP" The proportion of locations at which the genotype occurred in the top third (default).

Objects of class gai:

- "GAI" The geometric adaptability index (default).
- "GAI_R" The rank for the GAI values.

Objects of class superiority:

- "Pi_a" The superiority measure for all environments (default).
- "R_a" The rank for Pi_a.
- "Pi_f" The superiority measure for favorable environments.
- "R_f" The rank for Pi f.
- "Pi_u" The superiority measure for unfavorable environments.
- "R_u" The rank for Pi_u.

Objects of class Huehn:

- "S1" Mean of the absolute rank differences of a genotype over the n environments (default).
- "S2" variance among the ranks over the k environments.
- "S3" Sum of the absolute deviations.

- "S6" Relative sum of squares of rank for each genotype.
- "S1_R", "S2_R", "S3_R", and "S6_R", the ranks for S1, S2, S3, and S6, respectively.

Objects of class Thennarasu:

- "N1" First statistic (default).
- "N2" Second statistic.
- "N3" Third statistic.
- "N4" Fourth statistic.
- "N1_R", "N2_R", "N3_R", and "N4_R", The ranks for the statistics.

Objects of class performs_ammi:

- "PC1", "PC2", ..., "PCn" The values for the nth interaction principal component axis.
- "ipca_ss" Sum of square for each IPCA.
- "ipca_ms" Mean square for each IPCA.
- "ipca_fval" F value for each IPCA.
- "ipca_pval" P-value for for each IPCA.
- "ipca_expl" Explained sum of square for each IPCA (default).
- "ipca_accum" Accumulated explained sum of square.

Objects of class waas, waas_means, and waasb:

- "PC1", "PC2", ..., "PCn" The values for the nth interaction principal component axis.
- "WAASB" The weighted average of the absolute scores (default for objects of class waas).
- "PctResp" The rescaled values of the response variable.
- "PctWAASB" The rescaled values of the WAASB.
- "wResp" The weight for the response variable.
- "wWAASB" The weight for the stability.
- "OrResp" The ranking regarding the response variable.
- "OrWAASB" The ranking regarding the WAASB.
- "OrPC1" The ranking regarding the first principal component axix.
- "WAASBY" The superiority index WAASBY.
- "OrWAASBY" The ranking regarding the superiority index.

Objects of class gamem and waasb:

- "blupge" Best Linear Unbiased Prediction for genotype-environment interaction (mixed-effect model, class waasb).
- "blupg" Best Linear Unbiased Prediction for genotype effect.
- "bluege" Best Linear Unbiased Estimation for genotype-environment interaction (fixed-effect model, class waasb).
- "blueg" Best Linear Unbiased Estimation for genotype effect (fixed model).
- "data" The data used.
- "details" The details of the trial.
- "genpar" Genetic parameters (default).
- "gcov" The genotypic variance-covariance matrix.

- "pcov" The phenotypic variance-covariance matrix.
- "gcor" The genotypic correlation matrix.
- "pcor" The phenotypic correlation matrix.
- "h2" The broad-sense heritability.
- "lrt" The likelihood-ratio test for random effects.
- "vcomp" The variance components for random effects.
- "ranef" Random effects.

Objects of class blup_ind

- "HMGV", "HMGV_R" For harmonic mean of genotypic values or its ranks.
- "RPGV", RPGV_Y" For relative performance of genotypic values or its ranks.
- "HMRPGV", "HMRPGV_R" For harmonic mean of relative performance of genotypic values or its ranks.
- "WAASB", "WAASB_R" For the weighted average of absolute scores from the singular or its ranks. value decomposition of the BLUPs for GxE interaction or its ranks.

Value

A tibble showing the values of the variable chosen in argument what.

Author(s)

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See Also

```
AMMI_indexes(), anova_ind(), anova_joint(), ecovalence(), Fox(), gai(), gamem(), gafem(),
ge_acv(), ge_polar() ge_means(), ge_reg(), mgidi(), mtsi(), mps(), mtmps(), performs_ammi(),
blup_indexes(), Shukla(), superiority(), waas(), waasb()
```

```
library(metan)
ge_r <- ge_reg(data_ge2,</pre>
           env = ENV,
           gen = GEN,
           rep = REP,
           resp = c(PH, EH, CD, CL, ED))
get_model_data(ge_r)
# Significance of deviations from the regression
# Use gmd(), a shortcut for get_model_data
gmd(ge_r, "pval_f")
# Fitting the WAAS index
AMMI <- waasb(data_ge2,
           env = ENV,
           gen = GEN,
           rep = REP,
           resp = c(PH, ED, TKW, NKR))
# Getting the weighted average of absolute scores
gmd(AMMI, what = "WAASB")
# And the rank for the WAASB index.
gmd(AMMI, what = "OrWAASB")
# Fitting a mixed-effect model
# Genotype and interaction as random
blup <- gamem_met(data_ge2,</pre>
```

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ge_acv

Adjusted Coefficient of Variation as yield stability index

Description

[Stable]

Performs a stability analysis based on the scale-adjusted coefficient of variation (Doring and Reckling, 2018). For more details see acv()

Usage

```
ge_acv(.data, env, gen, resp, verbose = TRUE)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes and response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
resp	The response variable(s). To analyze multiple variables in a single procedure use, for example, $resp = c(var1, var2, var3)$.
verbose	Logical argument. If verbose = FALSE the code will run silently.

Value

An object of class ge_acv, which is a list containing the results for each variable used in the argument resp. For each variable, a tibble with the following columns is returned.

- **GEN** the genotype's code.
- ACV The adjusted coefficient of variation
- ACV_R The rank for the ACV value.

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Author(s)

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References

Doring, T.F., and M. Reckling. 2018. Detecting global trends of cereal yield stability by adjusting the coefficient of variation. Eur. J. Agron. 99: 30-36. doi: 10.1016/j.eja.2018.06.007

Examples

```
library(metan)
out <- ge_acv(data_ge2, ENV, GEN, c(EH, PH, EL, CD, ED, NKE))
gmd(out)</pre>
```

ge_cluster

Cluster genotypes or environments

Description

[Stable]

Performs clustering for genotypes or tester environments based on a dissimilarity matrix.

Usage

```
ge_cluster(
   .data,
   env = NULL,
   gen = NULL,
   table = FALSE,
   distmethod = "euclidean",
   clustmethod = "ward.D",
   scale = TRUE,
   cluster = "env",
   nclust = NULL
)
```

.data	The dataset containing the columns related to Environments, Genotypes and the response variable. It is also possible to use a two-way table with genotypes in lines and environments in columns as input. In this case you must use table = TRUE.
env	The name of the column that contains the levels of the environments. Defaults to NULL, in case of the input data is a two-way table.
gen	The name of the column that contains the levels of the genotypes. Defaults to NULL, in case of the input data is a two-way table.

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resp	The response variable(s). Defaults to NULL, in case of the input data is a two-way table.
table	Logical values indicating if the input data is a two-way table with genotypes in the rows and environments in the columns. Defaults to FALSE.
distmethod	The distance measure to be used. This must be one of 'euclidean', 'maximum', 'manhattan', 'canberra', 'binary', or 'minkowski'.
clustmethod	The agglomeration method to be used. This should be one of 'ward.D' (Default), 'ward.D2', 'single', 'complete', 'average' (= UPGMA), 'mcquitty (= WPGMA), 'median' (= WPGMC) or 'centroid' (= UPGMC).
scale	Should the data be scaled befor computing the distances? Set to TRUE. Let Y_{ij} be the yield of Hybrid i in Location j , $\bar{Y}_{.j}$ be the mean yield, and S_{j} be the standard deviation of Location j . The standardized yield (Zij) is computed as (Ouyang et al. 1995): $Z_{ij} = (Y_{ij} - Y_{.j})/S_{j}$.
cluster	What should be clustered? Defaults to cluster = "env" (cluster environments). To cluster the genotypes use cluster = "gen".
nclust	The number of clust to be formed. Set to NULL.

Value

- data The data that was used to compute the distances.
- **cutpoint** The cutpoint of the dendrogram according to Mojena (1977).
- distance The matrix with the distances.
- de The distances in an object of class dist.
- hc The hierarchical clustering.
- **cophenetic** The cophenetic correlation coefficient between distance matrix and cophenetic matrix
- Sqt The total sum of squares.
- tab A table with the clusters and similarity.
- **clusters** The sum of square and the mean of the clusters for each genotype (if cluster = "env" or environment (if cluster = "gen").
- labclust The labels of genotypes/environments within each cluster.

Author(s)

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References

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86 ge_details

Examples

```
library(metan)
d1 <- ge_cluster(data_ge, ENV, GEN, GY, nclust = 3)
plot(d1, nclust = 3)</pre>
```

ge_details

Details for genotype-environment trials

Description

[Stable]

Provide details for genotype-environment trials

Usage

```
ge_details(.data, env, gen, resp)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
resp	The response variable(s). To analyze multiple variables in a single procedure a vector of variables may be used. For example resp = c(var1, var2, var3). Select helpers are also allowed.

Value

A tibble with the following results for each variable:

- Mean: The grand mean.
- SE: The standard error of the mean.
- SD: The standard deviation.
- CV: The coefficient of variation.
- Min,Max: The minimum and maximum value, indicating the genotype and environment of occurence.
- MinENV, MinGEN: The environment and genotype with the lower mean.
- MaxENV, MaxGEN: The environment and genotype with the higher mean.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

ge_effects 87

Examples

```
library(metan)
details <- ge_details(data_ge2, ENV, GEN, everything())
print(details)</pre>
```

ge_effects

Genotype-environment effects

Description

[Stable]

This is a helper function that computes the genotype-environment effects, i.e., the residual effect of the additive model

Usage

```
ge_effects(.data, env, gen, resp, type = "ge", verbose = TRUE)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments. The analysis of variance is computed for each level of this factor.
gen	The name of the column that contains the levels of the genotypes.
resp	The response variable(s). To analyze multiple variables in a single procedure a vector of variables may be used. For example resp = c(var1, var2, var3).
type	The type of effect to compute. Defaults to "ge", i.e., genotype-environment. To compute genotype plus genotype-environment effects use type = "gge".
verbose	Logical argument. If verbose = FALSE the code will run silently.

Value

A list where each element is the result for one variable that contains a two-way table with genotypes in rows and environments in columns.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

```
library(metan)
ge_eff <- ge_effects(data_ge, ENV, GEN, GY)
gge_eff <- ge_effects(data_ge, ENV, GEN, GY, type = "gge")
plot(ge_eff)</pre>
```

88 ge_factanal

ge_factanal	Stability analysis and environment stratification	

Description

[Stable]

This function computes the stability analysis and environmental stratification using factor analysis as proposed by Murakami and Cruz (2004).

Usage

```
ge_factanal(.data, env, gen, rep, resp, mineval = 1, verbose = TRUE)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s)
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
rep	The name of the column that contains the levels of the replications/blocks
resp	The response variable(s). To analyze multiple variables in a single procedure use, for example, $resp = c(var1, var2, var3)$.
mineval	The minimum value so that an eigenvector is retained in the factor analysis.
verbose	Logical argument. If verbose = FALSE the code will run silently.

Value

An object of class ge_factanal with the following items:

- data: The data used to compute the factor analysis.
- cormat: The correlation matrix among the environments.
- PCA: The eigenvalues and explained variance.
- FA: The factor analysis.
- env_strat: The environmental stratification.
- KMO: The result for the Kaiser-Meyer-Olkin test.
- MSA: The measure of sampling adequacy for individual variable.
- communalities: The communalities.
- communalities.mean: The communalities' mean.
- initial.loadings: The initial loadings.
- finish.loadings: The final loadings after varimax rotation.
- canonical.loadings: The canonical loadings.
- scores.gen: The scores for genotypes for the first and second factors.

Author(s)

Tiago Olivoto, <tiagoolivoto@gmail.com>

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References

Murakami, D.M.D., and C.D.C. Cruz. 2004. Proposal of methodologies for environment stratification and analysis of genotype adaptability. Crop Breed. Appl. Biotechnol. 4:7-11.

See Also

```
superiority(), ecovalence(), ge_stats(), ge_reg()
```

Examples

ge_means

Genotype-environment means

Description

[Stable]

Computes genotype-environment interaction means

Usage

```
ge_means(.data, env, gen, resp)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, and the response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
resp	The response variable(s). To analyze multiple variables at once, a vector of variables may be used. For example $resp = c(var1, var2, var3)$. Select helpers are also allowed.

Value

A list where each element is the result for one variable containing:

- ge_means: A two-way table with the means for genotypes (rows) and environments (columns).
- gen_means: A tibble with the means for genotypes.
- env_means: A tibble with the means for environments.

90 ge_plot

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
means_ge <- ge_means(data_ge, ENV, GEN, resp = everything())
# Genotype-environment interaction means
get_model_data(means_ge)
# Environment means
get_model_data(means_ge, what = "env_means")
# Genotype means
get_model_data(means_ge, what = "gen_means")</pre>
```

ge_plot

Graphical analysis of genotype-vs-environment interaction

Description

[Stable]

This function produces a line plot for a graphical interpretation of the genotype-vs-environment interaction. By default, environments are in the x axis whereas the genotypes are depicted by different lines. The y axis contains the value of the selected variable. A heatmap can also be created.

Usage

```
ge_plot(
   .data,
   env,
   gen,
   resp,
   type = 1,
   plot_theme = theme_metan(),
   colour = TRUE
)
```

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments
gen	The name of the column that contains the levels of the genotypes.
resp	The response variable.

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type The type of plot type = 1 for a heatmap or type = 2 for a line plot.

plot_theme The graphical theme of the plot. Default is plot_theme = theme_metan(). For

more details, see ggplot2::theme().

colour Logical argument. If FALSE then the plot will not be colored.

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
ge_plot(data_ge2, ENV, GEN, PH)
ge_plot(data_ge, ENV, GEN, GY, type = 2)
```

ge_polar

Power Law Residuals as yield stability index

Description

[Stable]

Performs a stability analysis based on the Power Law Residuals (POLAR) statistics (Doring et al., 2015). POLAR is the residuals from the linear regression of $log(\sigma^2)$ against $log(\mu)$ and can be used as a measure of crop stability with lower stability (relative to all samples with that mean yield) indicated by more positive POLAR values, and higher stability (relative to all samples with that mean yield) indicated by more negative POLAR values.

Usage

```
ge_polar(.data, env, gen, resp, base = 10, verbose = TRUE)
```

.data	The dataset containing the columns related to Environments, Genotypes and response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
resp	The response variable(s). To analyze multiple variables in a single procedure use, for example, $resp = c(var1, var2, var3)$.
base	The base with respect to which logarithms are computed. Defaults to 10.
verbose	Logical argument. If verbose = FALSE the code will run silently.

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Value

An object of class ge_acv, which is a list containing the results for each variable used in the argument resp. For each variable, a tibble with the following columns is returned.

- **GEN** the genotype's code.
- POLAR The Power Law Residuals
- POLAR_R The rank for the ACV value.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

```
Doring, T.F., S. Knapp, and J.E. Cohen. 2015. Taylor's power law and the stability of crop yields. F. Crop. Res. 183: 294-302. doi: 10.1016/j.fcr.2015.08.005
```

Examples

```
library(metan)
out <- ge_polar(data_ge2, ENV, GEN, c(EH, PH, EL, CD, ED, NKE))
gmd(out)</pre>
```

ge_reg

Eberhart and Russell's regression model

Description

[Stable]

Regression-based stability analysis using the Eberhart and Russell (1966) model.

Usage

```
ge_reg(.data, env, gen, rep, resp, verbose = TRUE)
```

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response $variable(s)$
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
rep	The name of the column that contains the levels of the replications/blocks
resp	The response variable(s). To analyze multiple variables in a single procedure use, for example, $resp = c(var1, var2, var3)$.
verbose	Logical argument. If verbose = FALSE the code will run silently.

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Value

An object of class ge_reg with the folloing items for each variable:

 data: The data with means for genotype and environment combinations and the environment index

- anova: The analysis of variance for the regression model.
- regression: A data frame with the following columns: GEN, the genotypes; b0 and b1 the intercept and slope of the regression, respectively; t(b1=1) the calculated t-value; pval_t the p-value for the t test; s2di the deviations from the regression (stability parameter); F(s2di=0) the F-test for the deviations; pval_f the p-value for the F test; RMSE the root-mean-square error; R2 the determination coefficient of the regression.
- b0_variance: The variance of b0.
- b1_variance: The variance of b1.

Author(s)

Tiago Olivoto, <tiagoolivoto@gmail.com>

References

Eberhart, S.A., and W.A. Russell. 1966. Stability parameters for comparing Varieties. Crop Sci. 6:36-40. doi: 10.2135/cropsci1966.0011183X000600010011x

See Also

```
superiority(), ecovalence(), ge_stats()
```

Examples

ge_stats

Statistics for genotype-vs-environment interaction

Description

[Stable]

Computes (i) within-environment analysis of variance, GEI effect, GEI means, and genotype plus GEI effects; (ii) parametric statistics including AMMI-based indexes, Annicchiarico's genotypic confidence index (1992), Ecovalence (Wricke, 1965), regression-based stability (Eberhart and Russell., 1966), Shukla's stability variance parameter (1972); and (iii) nonparametric statistics including Fox's stability function (Fox et al. 1990), superiority index (Lin and Binns, 1988), Huehn's stability statistics (Huehn, 1979), and Thennarasu (1995) statistics.

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Usage

```
ge_stats(.data, env, gen, rep, resp, verbose = TRUE, prob = 0.05)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
rep	The name of the column that contains the levels of the replications/blocks.
resp	The response variable(s). To analyze multiple variables in a single procedure use, for example, resp = $c(var1, var2, var3)$.
verbose	Logical argument. If verbose = FALSE the code will run silently.
prob	The probability error assumed.

Details

The function computes the statistics and ranks for the following stability indexes. "Y" (Response variable), "CV" (coefficient of variation), "ACV" (adjusted coefficient of variation calling ge_acv() internally); POLAR (Power Law Residuals, calling ge_polar() internally) "Var" (Genotype's variance), "Shukla" (Shukla's variance, calling Shukla() internally), "Wi_g", "Wi_f", "Wi_u" (Annichiarrico's genotypic confidence index for all, favorable and unfavorable environments, respectively, calling Annicchiarico() internally), "Ecoval" (Wricke's ecovalence, ecovalence() internally), "Sij" (Deviations from the joint-regression analysis) and "R2" (R-squared from the jointregression analysis, calling ge_reg() internally), "ASV" (AMMI-stability value), "SIPC" (sum of the absolute values of the IPCA scores), "EV" (Average of the squared eigenvector values), "ZA" (Absolute values of the relative contributions of the IPCAs to the interaction), and "WAAS" (Weighted Average of Absolute Scores), by calling AMMI_indexes() internally; "HMGV" (Harmonic mean of the genotypic value), "RPGV" (Relative performance of the genotypic values), "HMRPGV" (Harmonic mean of the relative performance of the genotypic values), by calling blup_indexes() internally; "Pi_a", "Pi_f", "Pi_u" (Superiority indexes for all, favorable and unfavorable environments, respectively, calling superiority() internally), "Gai" (Geometric adaptability index, calling gai() internally), "S1" (mean of the absolute rank differences of a genotype over the n environments), "S2" (variance among the ranks over the k environments), "S3" (sum of the absolute deviations), "S6" (relative sum of squares of rank for each genotype), by calling Huehn() internally; and "N1", "N2", "N3", "N4" (Thennarasu"s statistics, calling Thennarasu() internally).

Value

An object of class ge_stats which is a list with one data frame for each variable containing the computed indexes.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

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Shukla, G.K. 1972. Some statistical aspects of partitioning genotype-environmental components of variability. Heredity. 29:238-245. doi: 10.1038/hdy.1972.87

Thennarasu, K. 1995. On certain nonparametric procedures for studying genotype x environment interactions and yield stability. Ph.D. thesis. P.J. School, IARI, New Delhi, India.

Wricke, G. 1965. Zur berechnung der okovalenz bei sommerweizen und hafer. Z. Pflanzenzuchtg 52:127-138.

Examples

```
library(metan)
model <- ge_stats(data_ge, ENV, GEN, REP, GY)
get_model_data(model, "stats")</pre>
```

ge_winners

Genotype-environment winners

Description

[Stable]

Computes the ranking for genotypes within environments and return the winners.

Usage

```
ge_winners(.data, env, gen, resp, type = "winners", better = NULL)
```

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Arguments

.data	The dataset containing the columns related to Environments, Genotypes, and the response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
resp	The response variable(s). To analyze multiple variables in a single procedure a vector of variables may be used. For example resp = c(var1, var2, var3). Select helpers are also allowed.
type	The type of results. Defaults to "winners" (default), i.e., a two-way table with the winner genotype in each environment. If type = "ranks" return the genotype ranking within each environment.
better	A vector of the same length of the number of variables to rank the genotypes according to the response variable. Each element of the vector must be one of the 'h' or 'l'. If 'h' is used (default), the genotypes are ranked from maximum to minimum. If 'l' is used then the are ranked from minimum to maximum. Use a comma-separated vector of names. For example, better = $c("h,h,h,h,l")$,

Value

A tibble with two-way table with the winner genotype in each environment (default) or the genotype ranking for each environment (if type = "ranks").

for ranking the fifth variable from minimum to maximum.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

gge 97

Description

[Stable]

Produces genotype plus genotype-by-environment model based on a multi-environment trial dataset containing at least the columns for genotypes, environments and one response variable or a two-way table.

Usage

```
gge(
   .data,
   env,
   gen,
   resp,
   centering = "environment",
   scaling = "none",
   svp = "environment",
   by = NULL,
   ...
)
```

Arguments

response variable(s).

env The name of the column that contains the levels of the environments.

gen The name of the column that contains the levels of the genotypes.

resp The response variable(s). To analyze multiple variables in a single procedure

a vector of variables may be used. For example resp = c(var1, var2, var3).

Select helpers are also supported.

centering The centering method. Must be one of the 'none | 0', for no centering; 'global

| 1', for global centered (E+G+GE); 'environment | 2' (default), for environment-centered (G+GE); or 'double | 3', for double centered (GE). A biplot cannot

be produced with models produced without centering.

scaling The scaling method. Must be one of the 'none | 0' (default), for no scaling;

or 'sd | 1', where each value is divided by the standard deviation of its corresponding environment (column). This will put all environments roughly he

same rang of values.

The method for singular value partitioning. Must be one of the 'genotype |

1', (The singular value is entirely partitioned into the genotype eigenvectors, also called row metric preserving); 'environment | 2', default, (The singular value is entirely partitioned into the environment eigenvectors, also called column metric preserving); or 'symmetrical | 3' (The singular value is symmetrically partitioned into the genotype and the environment eigenvectors This SVP is most often used in AMMI analysis and other biplot analysis, but it is not ideal for visualizing either the relationship among genotypes or that among the

environments).

by One variable (factor) to compute the function by. It is a shortcut to dplyr::group_by().This

is especially useful, for example, when the researcher want to produce GGE biplots for each level of a categorical variable. In this case, an object of class

gge_grouped is returned.

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... Arguments passed to the function impute_missing_val() for imputation of missing values in case of unbalanced data.

Value

The function returns a list of class gge containing the following objects

- coordgen The coordinates for genotypes for all components.
- coordenv The coordinates for environments for all components.
- eigenvalues The vector of eigenvalues.
- totalvar The overall variance.
- labelgen The name of the genotypes.
- labelenv The names of the environments.
- labelaxes The axes labels.
- ge_mat The data used to produce the model (scaled and centered).
- centering The centering method.
- scaling The scaling method.
- svp The singular value partitioning method.
- **d** The factor used to generate in which the ranges of genotypes and environments are comparable when singular value partitioning is set to 'genotype' or 'environment'.
- grand_mean The grand mean of the trial.
- mean_gen A vector with the means of the genotypes.
- mean_env A vector with the means of the environments.
- scale_var The scaling vector when the scaling method is 'sd'.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Yan, W., and M.S. Kang. 2003. GGE biplot analysis: a graphical tool for breeders, geneticists, and agronomists. CRC Press.

```
library(metan)
mod <- gge(data_ge, ENV, GEN, GY)
plot(mod)

# GGE model for all numeric variables
mod2 <- gge(data_ge2, ENV, GEN, resp = everything())
plot(mod2, var = "ED")

# If we have a two-way table with the mean values for
# genotypes and environments

table <- make_mat(data_ge, GEN, ENV, GY) %>% round(2)
table
make_long(table) %>%
gge(ENV, GEN, Y) %>%
plot()
```

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gtb Genotype by trait biplot	
------------------------------	--

Description

[Stable]

Produces a genotype-by-trait biplot model. From a genotype by environment by trait three-way table, genotype-by-trait tables in any single environment, across all environments, or across a subset of the environments can be generated and visually studied using biplots. The model for biplot analysis of genotype by trait data is the singular value decomposition of trait-standardized two-way table.

Usage

```
gtb(.data, gen, resp, centering = "trait", scaling = "sd", svp = "trait")
```

Arguments

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.data	The dataset containing the columns related to Genotypes and the response variable(s).
gen	The name of the column that contains the levels of the genotypes.
resp	The response variables, i.e., $resp = c(var1, var2, var3)$. Select helpers can also be used.
centering	The centering method. Must be one of the 'none $\mid \emptyset$ ', for no centering; 'global \mid 1', for global centered (T+G+GT); 'trait \mid 2' (default), for trait-centered (G+GT); or 'double \mid 3', for double centred (GT). A biplot cannot be produced with models produced without centering.
scaling	The scaling method. Must be one of the 'none 0', for no scaling; or 'sd 1' (default), where each value is divided by the standard deviation of its corresponding trait (column). This will put all traits roughly he same rang of values.
svp	The method for singular value partitioning. Must be one of the 'genotype 1', (The singular value is entirely partitioned into the genotype eigenvectors, also called row metric preserving); 'trait 2', default, (The singular value is entirely partitioned into the trait eigenvectors, also called column metric preserving); or 'symmetrical 3' (The singular value is symmetrically partitioned into the genotype and the trait eigenvectors This SVP is most often used in AMMI analysis and other biplot analysis, but it is not ideal for visualizing either the relationship among genotypes or that among the traits).

Value

The function returns a list of class gge that is compatible with the function plot() used in gge().

- coordgen The coordinates for genotypes for all components.
- coordenv The coordinates for traits for all components.
- eigenvalues The vector of eigenvalues.
- totalvar The overall variance.
- labelgen The name of the genotypes.

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- labelenv The names of the traits.
- labelaxes The axes labels.
- gt_mat The data used to produce the model (scaled and centered).
- **centering** The centering method.
- scaling The scaling method.
- svp The singular value partitioning method.
- **d** The factor used to generate in which the ranges of genotypes and traits are comparable when singular value partitioning is set to 'genotype' or 'trait'.
- grand_mean The grand mean of the trial.
- mean_gen A vector with the means of the genotypes.
- mean_env A vector with the means of the traits.
- scale_var The scaling vector when the scaling method is 'sd'.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Yan, W., and M.S. Kang. 2003. GGE biplot analysis: a graphical tool for breeders, geneticists, and agronomists. CRC Press.

Examples

```
library(metan)
# GT biplot for all numeric variables
mod <- gtb(data_ge2, GEN, resp = contains("E"))
plot(mod)</pre>
```

gytb

Genotype by yield*trait biplot

Description

[Stable]

Produces a Genotype by Yield*Trait biplot (GTY) proposed by Yan and Fregeau-Reid (2018).

Usage

```
gytb(
   .data,
   gen,
   yield,
   traits = everything(),
   ideotype = NULL,
   weight = NULL,
   prefix = "Y",
```

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```
centering = "trait",
scaling = "sd",
svp = "trait"
)
```

Arguments

.data The dataset containing the columns related to Genotypes, Yield, and Traits.

gen The name of the column that contains the levels of the genotypes.

yield The column containing the yield values.

traits The column(s) with the *traits* values. Defaults to *NULL*. In this case, all numeric

traits in .data, except that in yield are selected. To select specific traits from .data, use a list of unquoted comma-separated variable names (e.g. traits = c(var1, var2, var3)), an specific range of variables, (e.g. traits = c(var1:var3)),

or even a select helper like starts_with("N").

ideotype A vector of "h" or "l" with the same length of traits to define which trait is

desired to increase or decrease. By default (ideotype = NULL) for all numeric traits in traits are assumed that high values is desirable. Following the order of the traits selected in traits, use "h" to indicate the traits in which higher values are desired or "1" to indicate the variables in which lower values are desired. Then, yield will be multiplied by traits with "h" and divided by traits with "1" to generate the Genotype by yield*trait table. For example, ideotype = c("h,h,1") will assume that the ideotype has higher values for the first two

traits and lower values for the last trait.

weight The weight assumed for each trait. Similar to ideotype argument, provide a

numeric vector of the same length of traits. Suggested values are between $\boldsymbol{0}$

and 2.

prefix The prefix used in the biplot for the yield*trait combinations. Defaults to "Y".

centering The centering method. Must be one of the 'none | 0', for no centering; 'global

| 1', for global centered (T+G+GYT); 'trait | 2' (default), for trait-centered (G+GYT); or 'double | 3', for double centered (GYT). A biplot cannot be pro-

duced with models produced without centering.

scaling The scaling method. Must be one of the 'none | 0', for no scaling; or 'sd | 1'

(default), so that the mean for each trait or yield-trait combination becomes 0

and the variance becomes unit.

The method for singular value partitioning. Must be one of the 'genotype | 1',

(The singular value is entirely partitioned into the genotype eigenvectors, also called row metric preserving); 'trait | 2', default, (The singular value is entirely partitioned into the trait eigenvectors, also called column metric preserving); or 'symmetrical | 3' (The singular value is symmetrically partitioned into the genotype and the trait eigenvectors This SVP is most often used in AMMI analysis and other biplot analysis, but it is not ideal for visualizing either

the relationship among genotypes or that among the traits).

Value

The function returns a list of class gge that is compatible with the function plot() used in gge().

- data The Genotype by yield*trait (GYT) data.
- **ge_mat** The Genotype by yield*trait (GYT) data (scaled and centered).

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- coordgen The coordinates for genotypes for all components.
- coordenv The coordinates for traits for all components.
- eigenvalues The vector of eigenvalues.
- totalvar The overall variance.
- labelgen The name of the genotypes.
- labelenv The names of the traits.
- labelaxes The axes labels.
- **centering** The centering method.
- scaling The scaling method.
- svp The singular value partitioning method.
- **d** The factor used to generate in which the ranges of genotypes and traits are comparable when singular value partitioning is set to 'genotype' or 'trait'.
- grand_mean The grand mean of the trial.
- mean_gen A vector with the means of the genotypes.
- mean_env A vector with the means of the traits.
- scale_var The scaling vector when the scaling method is 'sd'.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Yan, W., & Fregeau-Reid, J. (2018). Genotype by Yield*Trait (GYT) Biplot: a Novel Approach for Genotype Selection based on Multiple Traits. Scientific Reports, 8(1), 1-10. doi: 10.1038/s41598018266888

Examples

```
library(metan)
# GYT biplot for all numeric traits of 'data_g'
# KW (kernel weight) considered as 'yield',
mod <- gytb(data_g, GEN, KW)
plot(mod)</pre>
```

Huehn

Huehn's stability statistics

Description

[Stable]

Performs a stability analysis based on Huehn (1979) statistics. The four nonparametric measures of phenotypic stability are: S1 (mean of the absolute rank differences of a genotype over the n environments), S2 (variance among the ranks over the k environments), S3 (sum of the absolute deviations), and S6 (relative sum of squares of rank for each genotype).

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Usage

```
Huehn(.data, env, gen, resp, verbose = TRUE)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
resp	The response variable(s). To analyze multiple variables in a single procedure use, for example, $resp = c(var1, var2, var3)$.
verbose	Logical argument. If verbose = FALSE the code will run silently.

Value

An object of class Huehn, which is a list containing the results for each variable used in the argument resp. For each variable, a tibble with the following columns is returned.

- **GEN** The genotype's code.
- Y The mean for the response variable.
- S1 Mean of the absolute rank differences of a genotype over the n environments.
- S2 variance among the ranks over the k environments.
- S3 Sum of the absolute deviations.
- S6 Relative sum of squares of rank for each genotype.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Huehn, V.M. 1979. Beitrage zur erfassung der phanotypischen stabilitat. EDV Med. Biol. 10:112.

```
library(metan)
out <- Huehn(data_ge2, ENV, GEN, PH)
print(out)</pre>
```

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impute_missing_val

Missing value imputation

Description

[Stable]

Impute the missing entries of a matrix with missing values using different algorithms. See **Details** section for more details

Usage

```
impute_missing_val(
   .data,
   naxis = 1,
   algorithm = "EM-SVD",
   tol = 1e-10,
   max_iter = 1000,
   simplified = FALSE,
   verbose = TRUE
)
```

Arguments

.data	A matrix to impute the missing entries. Frequently a two-way table of genotype means in each environment.
naxis	The rank of the Singular Value Approximation. Defaults to 1.
algorithm	The algorithm to impute missing values. Defaults to "EM-SVD". Other possible values are "EM-AMMI" and "colmeans". See Details section.
tol	The convergence tolerance for the algorithm.
max_iter	The maximum number of steps to take. If max_iter is achieved without convergence, the algorithm will stop with a warning.
simplified	Valid argument when algorithm = "EM-AMMI". IF FALSE (default), the current effects of rows and columns change from iteration to iteration. If TRUE, the general mean and effects of rows and columns are computed in the first iteration only, and in next iterations uses these values.
verbose	Logical argument. If verbose = FALSE the code will run silently.

Details

EM-AMMI algorithm

The EM-AMMI algorithm completes a data set with missing values according to both main and interaction effects. The algorithm works as follows (Gauch and Zobel, 1990):

- 1. The initial values are calculated as the grand mean increased by main effects of rows and main effects of columns. That way, the matrix of observations is pre-filled in.
- 2. The parameters of the AMMI model are estimated.
- 3. The adjusted means are calculated based on the AMMI model with naxis principal components.

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- 4. The missing cells are filled with the adjusted means.
- 5. The root mean square error of the predicted values (RMSE_p) is calculated with the two lasts iteration steps. If RMSE_p > tol, the steps 2 through 5 are repeated. Declare convergence if RMSE_p < tol. If max_iter is achieved without convergence, the algorithm will stop with a warning.

EM-SVD algorithm

The EM-SVD algorithm impute the missing entries using a low-rank Singular Value Decomposition approximation estimated by the Expectation-Maximization algorithm. The algorithm works as follows (Troyanskaya et al., 2001).

- 1. Initialize all NA values to the column means.
- 2. Compute the first naxis terms of the SVD of the completed matrix
- 3. Replace the previously missing values with their approximations from the SVD
- 4. The root mean square error of the predicted values (RMSE_p) is calculated with the two lasts iteration steps. If RMSE_p > tol, the steps 2 through 3 are repeated. Declare convergence if RMSE_p < tol. If max_iter is achieved without convergence, the algorithm will stop with a warning.

colmeans algorithm

The colmeans algorithm simply impute the missing entires using the column mean of the respective entire. Thus, there is no iteractive process.

Value

An object of class imv with the following values:

- .data The imputed matrix
- pc_ss The sum of squares representing variation explained by the principal components
- iter The final number of iterations.
- Final_RMSE The maximum change of the estimated values for missing cells in the last step of iteration.
- final_axis The final number of principal component axis.
- convergence Logical value indicating whether the modern converged.

References

Gauch, H. G., & Zobel, R. W. (1990). Imputing missing yield trial data. Theoretical and Applied Genetics, 79(6), 753-761. doi: 10.1007/BF00224240

Troyanskaya, O., Cantor, M., Sherlock, G., Brown, P., Hastie, T., Tibshirani, R., . Altman, R. B. (2001). Missing value estimation methods for DNA microarrays. Bioinformatics, 17(6), 520-525.

```
library(metan)
mat <- (1:20) %*% t(1:10)
mat
# 10% of missing values at random
miss_mat <- random_na(mat, prop = 10)
miss_mat
mod <- impute_missing_val(miss_mat)
mod$.data</pre>
```

inspect inspect

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	113	v		·	·

Check for common errors in multi-environment trial data

Description

[Stable]

inspect() scans a data.frame object for errors that may affect the use of functions in metan. By default, all variables are checked regarding the class (numeric or factor), missing values, and presence of possible outliers. The function will return a warning if the data looks like unbalanced, has missing values or possible outliers.

Usage

```
inspect(.data, ..., plot = FALSE, threshold = 15, verbose = TRUE)
```

Arguments

.data	The data to be analyzed
•••	The variables in .data to check. If no variable is informed, all the variables in .data are used.
plot	Create a plot to show the check? Defaults to FALSE.
threshold	Maximum number of levels allowed in a character / factor column to produce a plot. Defaults to 15.
verbose	Logical argument. If TRUE (default) then the results for checks are shown in the console.

Value

A tibble with the following variables:

- Variable The name of variable
- Class The class of the variable
- Missing Contains missing values?
- Levels The number of levels of a factor variable
- Valid_n Number of valid n (omit NAs)
- Outlier Contains possible outliers?

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

```
library(metan)
inspect(data_ge)

# Create a toy example with messy data
df <- data_ge2[-c(2, 30, 45, 134), c(1:5)]
df[c(1, 20, 50), c(4, 5)] <- NA</pre>
```

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```
df[40, 5] <- df[40, 5] * 2
inspect(df, plot = TRUE)</pre>
```

int.effects

Data for examples

Description

Data for examples

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

is.lpcor

Coerce to an object of class lpcor

Description

[Stable]

Functions to check if an object is of class 1pcor

Usage

```
is.lpcor(x)
```

Arguments

Χ

An object to check.

Value

A logical value TRUE or FALSE.

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is_balanced_trial	Check if a data set is balanced
13_batancca_trial	Check if a data set is bataneed

Description

Check if a data set coming from multi-environment trials is balanced, i.e., all genotypes are in all environments.

Usage

```
is_balanced_trial(.data, env, gen, resp)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
resp	The response variable.

Value

A logical value

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

lineplots

Fast way to create line plots

Description

[Stable]

- plot_lines() Creates a line plot based on one quantitative factor and one numeric variable. It can be used to show the results of a one-way trial with **quantitative treatments**.
- plot_factlines() Creates a line plot based on: one categorical and one quantitative factor and one numeric variable. It can be used to show the results of a two-way trial with qualitative-quantitative treatment structure.

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Usage

```
plot_lines(
  .data,
  Х,
  у,
  fit,
  level = 0.95,
  confidence = TRUE,
  xlab = NULL,
  ylab = NULL,
  n.dodge = 1,
  check.overlap = FALSE,
  col = "red",
  alpha = 0.2,
  size.shape = 1.5,
  size.line = 1,
  size.text = 12,
  fontfam = "sans",
  plot_theme = theme_metan()
)
plot_factlines(
  .data,
  Х,
  у,
  group,
  fit,
  level = 0.95,
  confidence = TRUE,
  xlab = NULL,
  ylab = NULL,
  n.dodge = 1,
  check.overlap = FALSE,
  legend.position = "bottom",
  grid = FALSE,
  scales = "free",
  col = TRUE,
  alpha = 0.2,
  size.shape = 1.5,
  size.line = 1,
  size.text = 12,
  fontfam = "sans",
  plot_theme = theme_metan()
)
```

Arguments

.data	The data set
x, y	The variables to be mapped to the x and y axes, respectively.
fit	The polynomial degree to use. It must be between 1 (linear fit) to 4 (fourth-order
	polynomial regression.). In plot_factlines(), if fit is a lenth 1 vector, i.e.,
	1, the fitted curves of all levels in group will be fitted with polynomial degree

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fit. To use a different polynomial degree for each level in group, use a numeric vector with the same length of the variable in group.

The fonfidence level. Defaults to 0.05.

confidence Display confidence interval around smooth? (TRUE by default) xlab, ylab The labels of the axes x and y, respectively. Defaults to NULL.

n. dodge The number of rows that should be used to render the x labels. This is useful for

displaying labels that would otherwise overlap.

check.overlap Silently remove overlapping labels, (recursively) prioritizing the first, last, and

middle labels.

col The colour to be used in the line plot and points.

alpha The alpha for the color in confidence band

size.shape The size for the shape in plot size.line The size for the line in the plot

size.text The size of the text

fontfam The family of the font text.

plot_theme The graphical theme of the plot. Default is plot_theme = theme_metan(). For

more details, see ggplot2::theme().

group The grouping variable. Valid for plot_factlines() only.

legend.position

level

Valid argument for plot_factlines. The position of the legend. Defaults to

'bottom'.

grid Valid argument for plot_factlines. Logical argument. If TRUE then a grid

will be created.

scales Valid argument for plot_factlines. If grid = TRUE scales controls how the

scales are in the plot. Possible values are 'free' (default), 'fixed', 'free_x'

or 'free_y'.

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

See Also

```
plot_bars() and plot_factbars()
```

Examples

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1pcor

Linear and Partial Correlation Coefficients

Description

[Stable]

Estimates the linear and partial correlation coefficients using as input a data frame or a correlation matrix.

Usage

```
lpcor(.data, ..., by = NULL, n = NULL, method = "pearson")
```

Arguments

.data	The data to be analyzed. It must be a symmetric correlation matrix or a data frame, possible with grouped data passed from dplyr::group_by().
	Variables to use in the correlation. If is null (Default) then all the numeric variables from .data are used. It must be a single variable name or a commaseparated list of unquoted variables names.
by	One variable (factor) to compute the function by. It is a shortcut to dplyr::group_by(). To compute the statistics by more than one grouping variable use that function.
n	If a correlation matrix is provided, then n is the number of objects used to compute the correlation coefficients.
method	a character string indicating which correlation coefficient is to be computed. One of 'pearson' (default), 'kendall', or 'spearman'.

Value

If .data is a grouped data passed from dplyr::group_by() then the results will be returned into a list-column of data frames, containing:

- linear.mat The matrix of linear correlation.
- partial.mat The matrix of partial correlations.
- results Hypothesis testing for each pairwise comparison.

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Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

mahala

Mahalanobis Distance

Description

[Stable]

Compute the Mahalanobis distance of all pairwise rows in .means. The result is a symmetric matrix containing the distances that may be used for hierarchical clustering.

Usage

```
mahala(.means, covar, inverted = FALSE)
```

Arguments

. means A matrix of data with, say, p columns.

covar The covariance matrix.

inverted Logical argument. If TRUE, covar is supposed to contain the inverse of the

covariance matrix.

Value

A symmetric matrix with the Mahalanobis' distance.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

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Examples

mahala_design

Mahalanobis distance from designed experiments

Description

[Stable]

Compute the Mahalanobis distance using data from an experiment conducted in a randomized complete block design or completely randomized design.

Usage

```
mahala_design(
   .data,
   gen,
   rep,
   resp,
   design = "RCBD",
   by = NULL,
   return = "distance"
)
```

Arguments

.data	The dataset containing the columns related to Genotypes, replication/block and response variables, possible with grouped data passed from dplyr::group_by().
gen	The name of the column that contains the levels of the genotypes.
rep	The name of the column that contains the levels of the replications/blocks.
resp	The response variables. For example $resp = c(var1, var2, var3)$.
design	The experimental design. Must be RCBD or CRD.

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by One variable (factor) to compute the function by. It is a shortcut to dplyr::group_by().

To compute the statistics by more than one grouping variable use that function.

return What the function return? Default is 'distance', i.e., the Mahalanobis distance.

Alternatively, it is possible to return the matrix of means return = 'means', or

the variance-covariance matrix of residuals return = 'covmat'.

Value

A symmetric matrix with the Mahalanobis' distance. If .data is a grouped data passed from dplyr::group_by() then the results will be returned into a list-column of data frames.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
maha <- mahala_design(data_g,</pre>
                       gen = GEN,
                       rep = REP,
                       resp = everything(),
                       return = "covmat")
# Compute one distance for each environment (all numeric variables)
maha_group <- mahala_design(data_ge,</pre>
                            gen = GEN,
                            rep = REP,
                            resp = everything(),
                            by = ENV)
# Return the variance-covariance matrix of residuals
cov_mat <- mahala_design(data_ge,</pre>
                         gen = GEN,
                         rep = REP,
                         resp = c(GY, HM),
                         return = 'covmat')
```

make_long

Two-way table to a 'long' format

Description

[Stable]

Helps users to easily convert a two-way table (genotype vs environment) to a 'long' format data. The data in mat will be gathered into three columns. The row names will compose the first column. The column names will compose the second column and the third column will contain the data that fills the two-way table.

Usage

```
make_long(mat, gen_in = "rows")
```

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Arguments

Mat A two-way table. It must be a matrix or a data.frame with rownames.

gen_in Where are the genotypes? Defaults to 'rows'. If genotypes are in columns and

environments in rows, set to gen_in = 'cols'.

Value

A tibble with three columns: GEN (genotype), ENV (environment), and Y (response) variable.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
set.seed(1)
mat <- matrix(rnorm(9, 2530, 350), ncol = 3)
colnames(mat) <- paste("E", 1:3, sep = "")
rownames(mat) <- paste("G", 1:3, sep = "")
make_long(mat)
gen_cols <- t(mat)
make_long(gen_cols, gen_in = "cols")</pre>
```

make_mat

Make a two-way table

Description

[Stable]

This function help users to easily make a two-way table from a "long format" data.

Usage

```
make_mat(.data, row, col, value, fun = mean)
```

Arguments

.data	The dataset. Must contains at least two categorical columns.
row	The column of data in which the mean of each level will correspond to one line in the output.
col	The column of data in which the mean of each level will correspond to one column in the output.
value	The column of data that contains the values to fill the two-way table.
fun	The function to apply. Defaults to mean, i.e., the two-way table will show the mean values for each genotype-environment combination. Other R base functions such as max, min, sd, var, or an own function that return a single numeric

value can be used.

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Value

A two-way table with the argument row in the rows, col in the columns, filled by the argument value.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
matrix <- data_ge %>% make_mat(row = GEN, col = ENV, val = GY)
matrix

# standart error of mean
data_ge %>% make_mat(GEN, ENV, GY, sem)
```

mantel_test

Mantel test

Description

[Stable]

Performs a Mantel test between two correlation/distance matrices. The function calculates the correlation between two matrices, the Z-score that is is the sum of the products of the corresponding elements of the matrices and a two-tailed p-value (null hypothesis:

r = 0

).

Usage

```
mantel_test(mat1, mat2, nboot = 1000, plot = FALSE)
```

Arguments

mat1, mat2 A correlation matrix or an object of class dist.

nboot The number of permutations to be used. Defaults to 1000.

plot if plot = TRUE, plots the density estimate of the permutation distribution along

with the observed Z-score as a vertical line.

Value

- mantel_r The correlation between the two matrices.
- z_score The Z-score.
- p-value The quantile of the observed Z-score. in the permutation distribution.

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Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

See Also

```
pairs_mantel()
```

Examples

```
library(metan)
# Test if the correlation of traits (data_ge2 dataset)
# changes between A1 and A2 levels of factor ENV
A1 <- corr_coef(data_ge2 %>% subset(ENV == "A1"))[["cor"]]
A2 <- corr_coef(data_ge2 %>% subset(ENV == "A2"))[["cor"]]
mantel_test(A1, A2, plot = TRUE)
```

meansGxE

Data for examples

Description

This dataset contains the means for grain yield of 10 genotypes cultivated in 5 environments. The interaction effects for this data is found in int.effects()

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

mgidi

Multitrait Genotype-Ideotype Distance Index

Description

[Stable]

Computes the multi-trait genotype-ideotype distance index, MGIDI, (Olivoto and Nardino, 2020), used to select genotypes in plant breeding programs based on multiple traits. The MGIDI index is computed as follows:

$$MGIDI_i = \sqrt{\sum_{j=1}^{f} (F_{ij} - F_j)^2}$$

where $MGIDI_i$ is the multi-trait genotype-ideotype distance index for the *i*th genotype; F_{ij} is the score of the *i*th genotype in the *j*th factor (i = 1, 2, ..., g; j = 1, 2, ..., f), being g and f the number of genotypes and factors, respectively, and F_j is the *j*th score of the ideotype. The genotype with the lowest MGIDI is then closer to the ideotype and therefore should presents desired values for all the analyzed traits.

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Usage

```
mgidi(
   .data,
   use_data = "blup",
   SI = 15,
   mineval = 1,
   ideotype = NULL,
   weights = NULL,
   use = "complete.obs",
   verbose = TRUE
)
```

Arguments

.data

An object fitted with the function gafem(), gamem() or a two-way table with BLUPs for genotypes in each trait (genotypes in rows and traits in columns). In the last case, row names must contain the genotypes names.

use_data

Define which data to use if .data is an object of class gamem. Defaults to "blup" (the BLUPs for genotypes). Use "pheno" to use phenotypic means instead BLUPs for computing the index.

SI

An integer (0-100). The selection intensity in percentage of the total number of genotypes.

mineval

The minimum value so that an eigenvector is retained in the factor analysis.

ideotype

A vector of length nvar where nvar is the number of variables used to plan the ideotype. Use 'h' to indicate the traits in which higher values are desired or 'l' to indicate the variables in which lower values are desired. For example, ideotype = c("h,h,h,h,l") will consider that the ideotype has higher values for the first four traits and lower values for the last trait. If .data is a model fitted with the functions gafem() or gamem(), the order of the traits will be the declared in the argument resp in those functions.

weights

Optional weights to assign for each trait in the selection process. It must be a numeric vector of length equal to the number of traits in .data. By default (NULL) a numeric vector of weights equal to 1 is used, i.e., all traits have the same weight in the selection process. It is suggested weights ranging from 0 to 1. The weights will then shrink the ideotype vector toward 0. This is useful, for example, to prioritize grain yield rather than a plant-related trait in the selection process.

use

The method for computing covariances in the presence of missing values. Defaults to complete.obs, i.e., missing values are handled by casewise deletion.

verbose

If verbose = TRUE (Default) then some results are shown in the console.

Value

An object of class mgidi with the following items:

- data The data used to compute the factor analysis.
- **cormat** The correlation matrix among the environments.
- **PCA** The eigenvalues and explained variance.
- FA The factor analysis.

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- KMO The result for the Kaiser-Meyer-Olkin test.
- MSA The measure of sampling adequacy for individual variable.
- communalities The communalities.
- communalities_mean The communalities' mean.
- initial_loadings The initial loadings.
- finish_loadings The final loadings after varimax rotation.
- canonical_loadings The canonical loadings.
- scores_gen The scores for genotypes in all retained factors.
- scores_ide The scores for the ideotype in all retained factors.
- gen ide The distance between the scores of each genotype with the ideotype.
- MGIDI The multi-trait genotype-ideotype distance index.
- **contri_fac** The relative contribution of each factor on the MGIDI value. The lower the contribution of a factor, the close of the ideotype the variables in such factor are.
- **contri_fac_rank, contri_fac_rank_sel** The rank for the contribution of each factor for all genotypes and selected genotypes, respectively.
- sel_dif The selection differential for the variables.
- **stat_gain** A descriptive statistic for the selection gains. The minimum, mean, confidence interval, standard deviation, maximum, and sum of selection gain values are computed. If traits have negative and positive desired gains, the statistics are computed for by strata.
- sel_gen The selected genotypes.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Olivoto, T., and Nardino, M. (2020). MGIDI: toward an effective multivariate selection in biological experiments. Bioinformatics. doi: 10.1093/bioinformatics/btaa981

Examples

```
library(metan)
# simulate a data set
# 10 genotypes
# 5 replications
# 4 traits
df <-
 g_simula(ngen = 10,
          nrep = 5,
          nvars = 4,
          gen_eff = 35,
          seed = c(1, 2, 3, 4))
# run a mixed-effect model (genotype as random effect)
mod <-
  gamem(df,
        gen = GEN,
        rep = REP,
```

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```
resp = everything())
# BLUPs for genotypes
gmd(mod, "blupg")
# Compute the MGIDI index
# Default options (all traits with positive desired gains)
# Equal weights for all traits
mgidi <- mgidi(mod)</pre>
gmd(mgidi, "MGIDI")
# Higher weight for traits V1 and V4
# This will increase the probability of selecting H7 and H9
# 30% selection pressure
mgidi2 <-
   mgidi(mod,
         weights = c(1, .2, .2, 1),
         SI = 30)
gmd(mgidi2, "MGIDI")
# plot the contribution of each factor on the MGIDI index
p1 <- plot(mgidi, type = "contribution")</pre>
p2 <- plot(mgidi2, type = "contribution")</pre>
p1 + p2
# Positive desired gains for V1, V2 and V3
# Negative desired gains for V4
mgidi3 <-
  mgidi(mod,
       ideotype = c("h, h, h, 1"))
```

mps

Mean performance and stability in multi-environment trials

Description

[Experimental]

This function implements the weighting method between mean performance and stability (Olivoto et al., 2019) considering different parametric and non-parametric stability indexes.

Usage

```
mps(
   .data,
   env,
   gen,
   rep,
   resp,
   block = NULL,
   by = NULL,
   random = "gen",
   performance = "blupg",
```

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```
stability = "waasb",
ideotype_mper = NULL,
ideotype_stab = NULL,
wmper = NULL,
verbose = TRUE
)
```

Arguments

cation/block and response variable(s).

env The name of the column that contains the levels of the environments.

gen The name of the column that contains the levels of the genotypes.

rep The name of the column that contains the levels of the replications/blocks.

resp The response variable(s). To analyze multiple variables in a single procedure a

vector of variables may be used. For example resp = c(var1, var2, var3).

block Defaults to NULL. In this case, a randomized complete block design is consid-

ered. If block is informed, then an alpha-lattice design is employed considering block as random to make use of inter-block information, whereas the complete replicate effect is always taken as fixed, as no inter-replicate information was to

be recovered (Mohring et al., 2015).

by One variable (factor) to compute the function by. It is a shortcut to dplyr::group_by().This

is especially useful, for example, when the researcher want to analyze environments within mega-environments. In this case, an object of class mps_grouped

is returned.

random The effects of the model assumed to be random. Defaults to random = "gen".

See gamem_met() to see the random effects assumed depending on the experi-

mental design of the trials.

performance Wich considers as mean performance. Either blupg (for Best Linear Unbiased

Prediction) or blueg (for Best Linear Unbiased Estimation)

stability The stability method. One of the following:

- "waasb" The weighted average of absolute scores (Olivoto et al. 2019).
- "ecovalence" The Wricke's ecovalence (Wricke, 1965).
- "Shukla" The Shukla's stability variance parameter (Shukla, 1972).
- "hmgv" The harmonic mean of genotypic values (Resende, 2007).
- "s2di" The deviations from the Eberhart and Russell regression (Eberhart and Russell, 1966).
- "r2" The determination coefficient of the Eberhart and Russell regression (Eberhart and Russell, 1966)...
- "rmse" The root mean squared error of the Eberhart and Russell regression (Eberhart and Russell, 1966).
- "wi" Annicchiarico's genotypic confidence index (Annicchiarico, 1992).
- "polar" Power Law Residuals as yield stability index (Doring et al., 2015).
- "acv" Adjusted Coefficient of Variation (Doring and Reckling, 2018)
- "pi" Lin e Binns' superiority index (Lin and Binns, 1988).
- "gai" Geometric adaptability index (Mohammadi and Amri, 2008).
- "s1", "s2", "s3", and "s6" Huehn's stability statistics (Huehn, 1979).

- "n1", "n2", "n3", and "n4" Thennarasu's stability statistics (Thennarasu, 1995).
- "asv", "ev", "za", and "waas" AMMI-based stability indexes (see AMMI_indexes()).

ideotype_mper, ideotype_stab

The new maximum value after rescaling the response variable/stability index. By default, all variables in resp are rescaled so that de maximum value is 100 and the minimum value is 0 (i.e., ideotype_mper = NULL and ideotype_stab = NULL). It must be a character vector of the same length of resp if rescaling is assumed to be different across variables, e.g., if for the first variable smaller values are better and for the second one, higher values are better, then ideotype_mper = c("1,h") must be used. For stability index in which lower values are better, use ideotype_stab = "1". Character value of length 1 will be recycled with a warning message.

wmper

The weight for the mean performance. By default, all variables in resp have equal weights for mean performance and stability (i.e., wmper = 50). It must be a numeric vector of the same length of resp to assign different weights across variables, e.g., if for the first variable equal weights for mean performance and stability are assumed and for the second one, a higher weight for mean performance (e.g. 65) is assumed, then wmper = c(50,65) must be used. Numeric value of length 1 will be recycled with a warning message.

verbose

Logical argument. If verbose = FALSE the code will run silently.

Value

An object of class mps with the following items.

- observed: The observed value on a genotype-mean basis.
- performance: The performance for genotypes (BLUPs or BLUEs)
- performance_res: The rescaled values of genotype's performance, considering ideotype_mper.
- stability: The stability for genotypes, chosen with argument stability.
- stability_res: The rescaled values of genotype's stability, considering ideotype_stab.
- mps_ind: The mean performance and stability for the traits.
- h2: The broad-sense heritability for the traits.
- perf_method: The method for measuring genotype's performance.
- wmper: The weight for the mean performance.
- sense_mper: The goal for genotype's performance (1 = lower, h = higher).
- stab_method: The method for measuring genotype's stability.
- wstab: The weight for the mean stability.
- sense_stab: The goal for genotype's stability (1 = lower, h = higher).

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

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See Also

```
mtsi(), mtmps(), mgidi()
```

Examples

```
library(metan)
# The same approach as mtsi()
# mean performance and stability for GY and HM
# mean performance: The genotype's BLUP
# stability: the WAASB index (lower is better)
# weights: equal for mean performance and stability

model <-
mps(data_ge,
    env = ENV,
    gen = GEN,
    rep = REP,
    resp = everything())

# The mean performance and stability after rescaling
model$mps_ind</pre>
```

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mtmps	Multi-trait mean performance and stability index
mtmps	Multi-trait mean performance and stability index

Description

[Experimental]

Computes the multi-trait stability index proposed by Olivoto et al. (2019) considering different parametric and non-parametric stability indexes.

Usage

```
mtmps(model, SI = 15, mineval = 1, verbose = TRUE)
```

Arguments

verbose

model An object of class mtmps

SI An integer (0-100). The selection intensity in percentage of the total number of genotypes.

mineval The minimum value so that an eigenvector is retained in the factor analysis.

If verbose = TRUE (Default), some results are shown in the console.

Value

An object of class mtmps with the following items:

- data The data used to compute the factor analysis.
- cormat The correlation matrix among the environments.
- PCA The eigenvalues and explained variance.
- FA The factor analysis.
- **KMO** The result for the Kaiser-Meyer-Olkin test.
- MSA The measure of sampling adequacy for individual variable.
- **communalities** The communalities.
- communalities_mean The communalities' mean.
- initial_loadings The initial loadings.
- finish loadings The final loadings after varimax rotation.
- canonical_loadings The canonical loadings.
- scores_gen The scores for genotypes in all retained factors.
- scores_ide The scores for the ideotype in all retained factors.
- MTSI The multi-trait mean performance and stability index.
- **contri_fac** The relative contribution of each factor on the MTSI value. The lower the contribution of a factor, the close of the ideotype the variables in such factor are.
- **contri_fac_rank, contri_fac_rank_sel** The rank for the contribution of each factor for all genotypes and selected genotypes, respectively.
- sel_dif_trait, sel_dif_stab, sel_dif_mps A data frame containing the selection differential (gains) for the mean performance, stability index, and mean performance and stability index, respectively. The following variables are shown.

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- VAR: the trait's name.
- Factor: The factor that traits where grouped into.
- Xo: The original population mean.
- Xs: The mean of selected genotypes.
- SD and SDperc: The selection differential and selection differential in percentage, respectively.
- h2: The broad-sense heritability.
- SG and SGperc: The selection gains and selection gains in percentage, respectively.
- sense: The desired selection sense.
- goal: selection gains match desired sense? 100 for yes and 0 for no.
- stat_dif_trait, stat_dif_stab, stat_dif_mps A data frame with the descriptive statistic for the selection gains for the mean performance, stability index, and mean performance and stability index, respectively. The following columns are shown by sense.
 - sense: The desired selection sense.
 - variable: the trait's name.
 - min: the minimum value for the selection gain.
 - mean: the mean value for the selection gain.
 - ci: the confidence interval for the selection gain.
 - sd. amo: the standard deviation for the selection gain.
 - max: the maximum value for the selection gain.
 - sum: the sum of the selection gain.
- sel_gen The selected genotypes.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Olivoto, T., A.D.C. L\'ucio, J.A.G. da silva, B.G. Sari, and M.I. Diel. 2019. Mean performance and stability in multi-environment trials II: Selection based on multiple traits. Agron. J. 111:2961-2969. doi: 10.2134/agronj2019.03.0220

See Also

```
mgidi(), mps(), get_model_data()
```

Examples

```
library(metan)
# The same approach as mtsi()
# mean performance and stability for GY and HM
# mean performance: The genotype's BLUP
# stability: the WAASB index (lower is better)
# weights: equal for mean performance and stability

model <-
mps(data_ge,
    env = ENV,
    gen = GEN,
    rep = REP,</pre>
```

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```
resp = everything())
selection <- mtmps(model)

# gains for stability
selection$sel_dif_stab

# gains for mean performance
selection$sel_dif_trait</pre>
```

mtsi

Multi-trait stability index

Description

[Stable]

Computes the multi-trait stability index proposed by Olivoto et al. (2019)

Usage

```
mtsi(
   .data,
   index = "waasby",
   ideotype = NULL,
   SI = 15,
   mineval = 1,
   verbose = TRUE
)
```

Arguments

	.data	An object of class v	waasb or waas.
--	-------	----------------------	----------------

index If index = 'waasby' (default) both stability and mean performance are considered. If index = 'waasb' the multi-trait index will be computed considering the

ered. If index = 'waasb' the multi-trait index will be computed considering the stability of genotypes only. More details can be seen in waasb() and waas()

functions.

ideotype A vector of length nvar where nvar is the number of variables used to plan the

ideotype. Use 'h' to indicate the traits in which higher values are desired or '1' to indicate the variables in which lower values are desired. For example, ideotype = c("h,h,h,h,h,1") will consider that the ideotype has higher values

for the first four traits and lower values for the last trait.

SI An integer (0-100). The selection intensity in percentage of the total number of

genotypes.

mineval The minimum value so that an eigenvector is retained in the factor analysis.

verbose If verbose = TRUE (Default) then some results are shown in the console.

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Value

An object of class mtsi with the following items:

- data The data used to compute the factor analysis.
- **cormat** The correlation matrix among the environments.
- PCA The eigenvalues and explained variance.
- **FA** The factor analysis.
- KMO The result for the Kaiser-Meyer-Olkin test.
- MSA The measure of sampling adequacy for individual variable.
- communalities The communalities.
- communalities_mean The communalities' mean.
- initial_loadings The initial loadings.
- finish_loadings The final loadings after varimax rotation.
- canonical_loadings The canonical loadings.
- scores_gen The scores for genotypes in all retained factors.
- scores_ide The scores for the ideotype in all retained factors.
- MTSI The multi-trait stability index.
- **contri_fac** The relative contribution of each factor on the MTSI value. The lower the contribution of a factor, the close of the ideotype the variables in such factor are.
- **contri_fac_rank, contri_fac_rank_sel** The rank for the contribution of each factor for all genotypes and selected genotypes, respectively.
- sel_dif_trait, sel_dif_stab, sel_dif_mps A data frame containing the selection differential (gains) for the traits, for the stability (WAASB index) WAASB, and for the mean performance and stability (WAASBY indexes). The following variables are shown.
 - VAR: the trait's name.
 - Factor: The factor that traits where grouped into.
 - Xo: The original population mean.
 - Xs: The mean of selected genotypes.
 - SD and SDperc: The selection differential and selection differential in percentage, respectively.
 - h2: The broad-sense heritability.
 - SG and SGperc: The selection gains and selection gains in percentage, respectively.
 - sense: The desired selection sense.
 - goal: selection gains match desired sense? 100 for yes and 0 for no.
- stat_dif_var, stat_dif_stab, stat_dif_mps A data frame with the descriptive statistic for the selection gains for the traits, for the stability (WAASB index) WAASB, and for the mean performance and stability (WAASBY index). The following variables are shown.
 - sense: The desired selection sense.
 - variable: the trait's name.
 - min: the minimum value for the selection gain.
 - mean: the mean value for the selection gain.
 - ci: the confidence interval for the selection gain.
 - sd. amo: the standard deviation for the selection gain.
 - max: the maximum value for the selection gain.
 - sum: the sum of the selection gain.
- sel_gen The selected genotypes.

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Author(s)

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References

Olivoto, T., A.D.C. L\'ucio, J.A.G. da silva, B.G. Sari, and M.I. Diel. 2019. Mean performance and stability in multi-environment trials II: Selection based on multiple traits. Agron. J. 111:2961-2969. doi: 10.2134/agronj2019.03.0220

See Also

```
mgidi(), waasb(), get_model_data()
```

Examples

```
library(metan)
# Based on stability only, for both GY and HM, higher is better
mtsi_model <-
    waasb(data_ge,
    env = ENV,
    gen = GEN,
    rep = REP,
    resp = c(GY, HM))
mtsi_index <-</pre>
    mtsi(mtsi_model, index = 'waasb')
# Based on mean performance and stability (using pipe operator %>%)
# GY: higher is better
# HM: lower is better
mtsi_index2 <-</pre>
data_ge %>%
 waasb(ENV, GEN, REP,
       resp = c(GY, HM),
       mresp = c("h, 1")) \%>\%
 mtsi()
```

non_collinear_vars

Select a set of predictors with minimal multicollinearity

Description

[Stable]

Select a set of predictors with minimal multicollinearity using the variance inflation factor (VIF) as criteria to remove collinear variables. The algorithm will: (i) compute the VIF value of the correlation matrix containing the variables selected in ...; (ii) arrange the VIF values and delete the variable with the highest VIF; and (iii) iterate step ii until VIF value is less than or equal to max_vif.

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Usage

```
non_collinear_vars(
   .data,
   ...,
   max_vif = 10,
   missingval = "pairwise.complete.obs"
)
```

Arguments

.data	The data set containing the variables.
	Variables to be submitted to selection. If is null then all the numeric variables from .data are used. It must be a single variable name or a commaseparated list of unquoted variables names.
max_vif	The maximum value for the Variance Inflation Factor (threshold) that will be accepted in the set of selected predictors.
missingval	How to deal with missing values. For more information, please see stats::cor().

Value

A data frame showing the number of selected predictors, maximum VIF value, condition number, determinant value, selected predictors and removed predictors from the original set of variables.

Examples

```
library(metan)
# All numeric variables
non_collinear_vars(data_ge2)

# Select variables and choose a VIF threshold to 5
non_collinear_vars(data_ge2, EH, CL, CW, KW, NKE, max_vif = 5)
```

pairs_mantel

Mantel test for a set of correlation matrices

Description

[Stable]

This function generate a pairwise matrix of plots to compare the similarity of two or more correlation matrices. In the upper diagonal are presented the plots and in the lower diagonal the result of Mantel test based on permutations.

Usage

```
pairs_mantel(
    ...,
    type = 1,
    nrepet = 1000,
    names = NULL,
```

pairs_mantel

```
prob = 0.05,
  diag = FALSE,
  export = FALSE,
  main = "auto",
  file.type = "pdf",
  file.name = NULL,
  width = 8,
  height = 7,
  resolution = 300,
  size.point = 0.5,
  shape.point = 19,
  alpha.point = 1,
  fill.point = NULL,
  col.point = "black",
  minsize = 2,
 maxsize = 3,
  signcol = "green",
  alpha = 0.15,
  diagcol = "gray",
  col.up.panel = "gray",
  col.lw.panel = "gray",
  col.dia.panel = "gray",
  pan.spacing = 0.15,
  digits = 2
)
```

Arguments

fill.point

	The input matrices. May be an output generated by the function 1pcor or a coerced list generated by the function as.1pcor
type	The type of correlation if an obect generated by the function 1pcor is used. $1 =$ Linear correlation matrices, or $2 =$ partial correlation matrices.
nrepet	The number of permutations. Default is 1000
names	An optional vector of names of the same length of
prob	The error probability for Mantel test.
diag	Logical argument. If TRUE, the Kernel density is shown in the diagonal of plot.
export	Logical argument. If TRUE, then the plot is exported to the current directory.
main	The title of the plot, set to 'auto'.
file.type	The format of the file if export = TRUE. Set to 'pdf'. Other possible values are *.tiff using file.type = 'tiff'.
file.name	The name of the plot when exported. Set to NULL, i.e., automatically.
width	The width of the plot, set to 8.
height	The height of the plot, set to 7.
resolution	The resolution of the plot if file.type = 'tiff' is used. Set to 300 (300 dpi).
size.point	The size of the points in the plot. Set to 0.5 .
shape.point	The shape of the point, set to 19.
alpha.point	The value for transparency of the points: $1 = \text{full color}$.

The color to fill the points. Valid argument if points are between 21 and 25.

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col.point	The color for the edge of the point, set to black.
minsize	The size of the letter that will represent the smallest correlation coefficient.
maxsize	The size of the letter that will represent the largest correlation coefficient.
signcol	The colour that indicate significant correlations (based on the prob value.), set to 'green'.
alpha	The value for transparency of the color informed in signcol, when $1 = \text{full}$ color. Set to 0.15 .
diagcol	The color in the kernel distribution. Set to 'gray'.
col.up.panel, c	ol.lw.panel, col.dia.panel The color for the opper, lower and diagonal pannels. Set to 'gray', 'gray', and 'gray', respectively.
pan.spacing	The space between the pannels. Set to 0.15.
digits	The number of digits to show in the plot.

Value

An object of class gg, ggmatrix.

Author(s)

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See Also

```
mantel_test()
```

Examples

path_coeff

path_coeff

Path coefficients with minimal multicollinearity

Description

[Stable]

- path_coeff() computes a path analysis using a data frame as input data.
- path_coeff_mat() computes a path analysis using correlation matrices as input data.

Usage

```
path_coeff(
    .data,
    resp,
    pred = everything(),
    by = NULL,
    exclude = FALSE,
    correction = NULL,
    knumber = 50,
    brutstep = FALSE,
    maxvif = 10,
    missingval = "pairwise.complete.obs",
    plot_res = FALSE,
    verbose = TRUE,
    ...
)

path_coeff_mat(cor_mat, resp, correction = NULL, knumber = 50, verbose = TRUE)
```

Arguments

.data	The data. Must be a data frame or a grouped data passed from dplyr::group_by()
resp	The dependent variable.
pred	The predictor variables, set to everything(), i.e., the predictor variables are all the numeric variables in the data except that in resp.
by	One variable (factor) to compute the function by. It is a shortcut to dplyr::group_by(). To compute the statistics by more than one grouping variable use that function.
exclude	Logical argument, set to false. If exclude = TRUE, then the variables in pred are deleted from the data, and the analysis will use as predictor those that remained, except that in resp.
correction	Set to NULL. A correction value (k) that will be added into the diagonal elements of the X'X matrix aiming at reducing the harmful problems of the multicollinearity in path analysis (Olivoto et al., 2017)
knumber	When correction = NULL, a plot showing the values of direct effects in a set of different k values (0-1) is produced. knumber is the number of k values used in the range of 0 to 1.
brutstep	Logical argument, set to FALSE. If true, then an algorithm will select a subset

the **Details** section for more information.

of variables with minimal multicollinearity and fit a set of possible models. See

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maxvif	The maximum value for the Variance Inflation Factor (cut point) that will be accepted. See the Details section for more information.
missingval	How to deal with missing values. For more information, please see stats::cor().
plot_res	If TRUE, create a scatter plot of residual against predicted value and a normal Q-Q plot.
verbose	If verbose = TRUE then some results are shown in the console.
	Additional arguments passed on to stats::plot.lm()
cor_mat	Matrix of correlations containing both dependent and independent traits.

Details

In path_coeff(), when brutstep = TRUE, an algorithm to select a set of predictors with minimal multicollinearity and high explanatory power is implemented. first, the algorithm will select a set of predictors with minimal multicollinearity. The selection is based on the variance inflation factor (VIF). An iterative process is performed until the maximum VIF observed is less than maxvif. The variables selected in this iterative process are then used in a series of stepwise-based regressions. The first model is fitted and p-1 predictor variables are retained (p is the number of variables selected in the iterative process. The second model adjusts a regression considering p-2 selected variables, and so on until the last model, which considers only two variables. Three objects are created. Summary, with the process summary, Models, containing the aforementioned values for all the adjusted models; and Selectedpred, a vector with the name of the selected variables in the iterative process.

Value

An object of class path_coeff, group_path, or brute_path with the following items:

- Corr.x A correlation matrix between the predictor variables.
- Corr.y A vector of correlations between each predictor variable with the dependent variable.
- Coefficients The path coefficients. Direct effects are the diagonal elements, and the indirect effects those in the off-diagonal elements (lines).
- **Eigen** Eigenvectors and eigenvalues of the Corr.x.
- VIF The Variance Inflation Factors.
- plot A ggplot2-based graphic showing the direct effects in 21 different k values.
- **Predictors** The predictor variables used in the model.
- CN The Condition Number, i.e., the ratio between the highest and lowest eigenvalue.
- **Det** The matrix determinant of the Corr.x..
- **R2** The coefficient of determination of the model.
- Residual The residual effect of the model.
- **Response** The response variable.
- **weightvar** The order of the predictor variables with the highest weight (highest eigenvector) in the lowest eigenvalue.

If .data is a grouped data passed from dplyr::group_by() then the results will be returned into a list-column of data frames, containing:

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

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References

Olivoto, T., V.Q. Souza, M. Nardino, I.R. Carvalho, M. Ferrari, A.J. Pelegrin, V.J. Szareski, and D. Schmidt. 2017. Multicollinearity in path analysis: a simple method to reduce its effects. Agron. J. 109:131-142. doi: 10.2134/agronj2016.04.0196

Examples

```
library(metan)
# Using KW as the response variable and all other ones as predictors
pcoeff <- path_coeff(data_ge2, resp = KW)</pre>
# The same as above, but using the correlation matrix
cor_mat <- cor(data_ge2 %>% select_numeric_cols())
pcoeff <- path_coeff_mat(cor_mat, resp = KW)</pre>
# Declaring the predictors
# Create a residual plot with 'plot_res = TRUE'
pcoeff2 <- path_coeff(data_ge2,</pre>
                       resp = KW,
                       pred = c(PH, EH, NKE, TKW),
                       plot_res = TRUE)
# Selecting variables to be excluded from the analysis
pcoeff3 <- path_coeff(data_ge2,</pre>
                      resp = KW,
                      pred = c(NKR, PERK, KW, NKE),
                      exclude = TRUE)
# Selecting a set of predictors with minimal multicollinearity
# Maximum variance Inflation factor of 5
pcoeff4 <- path_coeff(data_ge2,</pre>
                      resp = KW.
                      brutstep = TRUE,
                      maxvif = 5)
# When one analysis should be carried out for each environment
# Using the forward-pipe operator %>%
pcoeff5 <- path_coeff(data_ge2, resp = KW, by = ENV)</pre>
```

performs_ammi

Additive Main effects and Multiplicative Interaction

Description

[Stable]

Compute the Additive Main effects and Multiplicative interaction (AMMI) model. The estimate of the response variable for the *i*th genotype in the *j*th environment (y_{ij}) using the AMMI model, is given as follows:

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$$y_{ij} = \mu + \alpha_i + \tau_j + \sum_{k=1}^p \lambda_k a_{ik} t_{jk} + \rho_{ij} + \varepsilon_{ij}$$

where λ_k is the singular value for the k-th interaction principal component axis (IPCA); a_{ik} is the i-th element of the k-th eigenvector; t_{jk} is the jth element of the kth eigenvector. A residual ρ_{ij} remains, if not all p IPCA are used, where $p \leq min(g-1;e-1)$.

This function also serves as a helper function for other procedures performed in the **metan** package such as waas() and wsmp()

Usage

```
performs_ammi(.data, env, gen, rep, resp, block = NULL, verbose = TRUE, ...)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments
gen	The name of the column that contains the levels of the genotypes
rep	The name of the column that contains the levels of the replications/blocks
resp	The response variable(s). To analyze multiple variables in a single procedure, use comma-separated list of unquoted variable names, i.e., resp = c(var1, var2, var3), or any select helper like resp = contains("_PLA").
block	Defaults to NULL. In this case, a randomized complete block design is considered. If block is informed, then a resolvable alpha-lattice design (Patterson and Williams, 1976) is employed. All effects, except the error, are assumed to be fixed.
verbose	Logical argument. If verbose = FALSE the code will run silently.
	Arguments passed to the function impute_missing_val() for imputation of missing values in case of unbalanced data.

Value

- ANOVA: The analysis of variance for the AMMI model.
- PCA: The principal component analysis
- MeansGxE: The means of genotypes in the environments
- model: scores for genotypes and environments in all the possible axes.
- augment: Information about each observation in the dataset. This includes predicted values in the fitted column, residuals in the resid column, standardized residuals in the stdres column, the diagonal of the 'hat' matrix in the hat, and standard errors for the fitted values in the se.fit column.

Author(s)

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plot.anova_joint

References

Patterson, H.D., and E.R. Williams. 1976. A new class of resolvable incomplete block designs. Biometrika 63:83-92.

See Also

```
impute_missing_val(), waas(), waas_means(), waasb(), get_model_data()
```

Examples

```
library(metan)
model <- performs_ammi(data_ge, ENV, GEN, REP, resp = c(GY, HM))</pre>
# PC1 x PC2 (variable GY)
p1 <- plot_scores(model)</pre>
p1
# PC1 x PC2 (variable HM)
plot_scores(model,
            var = 2, # or "HM"
            type = 2)
# Nominal yield plot (variable GY)
# Draw a convex hull polygon
plot_scores(model, type = 4)
# Unbalanced data (GEN 2 in E1 missing)
mod <-
  data_ge %>%
   remove_rows(4:6) %>%
   droplevels() %>%
   performs_ammi(ENV, GEN, REP, GY)
p2 <- plot_scores(mod)</pre>
arrange_ggplot(p1, p2, tag_levels = list(c("Balanced data", "Unbalanced data")))
```

plot.anova_joint

Several types of residual plots

Description

Residual plots for a output model of class anova_joint. Seven types of plots are produced: (1) Residuals vs fitted, (2) normal Q-Q plot for the residuals, (3) scale-location plot (standardized residuals vs Fitted Values), (4) standardized residuals vs Factor-levels, (5) Histogram of raw residuals and (6) standardized residuals vs observation order, and (7) 1:1 line plot.

Usage

```
## S3 method for class 'anova_joint'
plot(x, ...)
```

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Arguments

```
x An object of class anova_joint.... Additional arguments passed on to the function residual_plots()
```

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
model <- anova_joint(data_ge, ENV, GEN, REP, GY)
plot(model)
plot(model,
    which = c(3, 5),
    nrow = 2,
    labels = TRUE,
    size.lab.out = 4)</pre>
```

plot.can_cor

Plots an object of class can_cor

Description

Graphs of the Canonical Correlation Analysis

Usage

```
## S3 method for class 'can_cor'
plot(
  х,
  type = 1,
  plot_theme = theme_metan(),
  size.tex.lab = 12,
  size.tex.pa = 3.5,
  x.lab = NULL,
  x.lim = NULL,
  x.breaks = waiver(),
  y.lab = NULL,
  y.lim = NULL,
  y.breaks = waiver(),
  axis.expand = 1.1,
  shape = 21,
  col.shape = "orange",
  col.alpha = 0.9,
  size.shape = 3.5,
  size.bor.tick = 0.3,
  labels = FALSE,
  main = NULL,
)
```

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Arguments

x	The waasb object
type	The type of the plot. Defaults to type = 1 (Scree-plot of the correlations of the canonical loadings). Use type = 2, to produce a plot with the scores of the variables in the first group, type = 3 to produce a plot with the scores of the variables in the second group, or type = 4 to produce a circle of correlations.
plot_theme	The graphical theme of the plot. Default is plot_theme = theme_metan(). For more details,see ggplot2::theme().
size.tex.lab	The size of the text in axis text and labels.
size.tex.pa	The size of the text of the plot area. Default is 3.5.
x.lab	The label of x-axis. Each plot has a default value. New arguments can be inserted as $x.lab = 'my label'$.
x.lim	The range of x-axis. Default is NULL (maximum and minimum values of the data set). New arguments can be inserted as $x.lim = c(x.min, x.max)$.
x.breaks	The breaks to be plotted in the x-axis. Default is authomatic breaks. New arguments can be inserted as x . breaks = $c(breaks)$
y.lab	The label of y-axis. Each plot has a default value. New arguments can be inserted as y.lab = 'my label'.
y.lim	The range of y-axis. Default is NULL. The same arguments than $x.lim$ can be used.
y.breaks	The breaks to be plotted in the x-axis. Default is authomatic breaks. The same arguments than x.breaks can be used.
axis.expand	Multiplication factor to expand the axis limits by to enable fitting of labels. Default is 1.1.
shape	The shape of points in the plot. Default is 21 (circle). Values must be between 21-25: 21 (circle), 22 (square), 23 (diamond), 24 (up triangle), and 25 (low triangle).
col.shape	A vector of length 2 that contains the color of shapes for genotypes above and below of the mean, respectively. Defaults to "orange". c("blue", "red").
col.alpha	The alpha value for the color. Default is 0.9 . Values must be between 0 (full transparency) to 1 (full color).
size.shape	The size of the shape in the plot. Default is 3.5.
size.bor.tick	The size of tick of shape. Default is 0.3. The size of the shape will be size.shape $+$ size.bor.tick
labels	Logical arguments. If TRUE then the points in the plot will have labels.
main	The title of the plot. Defaults to NULL, in which each plot will have a default title. Use a string text to create an own title or set to main = FALSE to omit the plot title.
• • •	Currently not used.

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

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Examples

plot.clustering

Plot an object of class clustering

Description

Plot an object of class clustering

Usage

```
## S3 method for class 'clustering'
plot(x, horiz = TRUE, type = "dendrogram", ...)
```

Arguments

Х	An object of class clustering
horiz	Logical indicating if the dendrogram should be drawn horizontally or not.
type	The type of plot. Must be one of the 'dendrogram' or 'cophenetic'.
	Other arguments passed from the function plot.dendrogram or abline.

Value

An object of class gg, ggplot if type == "cophenetic".

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

plot.corr_coef

Examples

```
mean_gen <-
  data_ge2 %>%
  means_by(GEN) %>%
  column_to_rownames("GEN")

d <- clustering(mean_gen)
plot(d, xlab = "Euclidean Distance")</pre>
```

plot.correlated_vars
Plot an object of class correlated_vars

Description

Plot an object of class correlated_vars

Usage

```
## S3 method for class 'correlated_vars'
plot(x, ...)
```

Arguments

x An object of class correlated_vars.

... Currently not used.

Value

An object of class gg.

Examples

```
library(metan)
y <- rnorm(n = 10)
cor_vars <- correlated_vars(y, nvar = 6)
plot(cor_vars)</pre>
```

plot.corr_coef

Create a correlation heat map

Description

Create a correlation heat map for object of class corr_coef

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Usage

```
## S3 method for class 'corr_coef'
plot(
  Х,
  type = "lower",
  diag = FALSE,
  reorder = TRUE,
  signif = "stars",
  caption = TRUE,
  digits.cor = 2,
  digits.pval = 3,
  col.low = "blue",
  col.mid = "white",
  col.high = "red",
  lab.x.position = NULL,
  lab.y.position = NULL,
  legend.position = NULL,
  legend.title = "Pearson's\nCorrelation",
  size.text.cor = 3,
  size.text.signif = 3,
  size.text.lab = 10,
```

Arguments

The data set. The type of heat map to produce. Either lower (default) to produce a lower type triangle heat map or upper to produce an upper triangular heat map. diag Plot diagonal elements? Defaults to FALSE. reorder Reorder the correlation matrix to identify the hidden pattern? Defaults to TRUE. signif How to show significant correlations. If "stars" is used (default), stars are used showing the significance at 0.05 (""), 0.01 ("") and 0.001 ("") probability error. If signif = "pval", then the p-values are shown. Logical. If TRUE (Default) includes a caption with the significance meaning for caption stars. digits.cor, digits.pval The significant digits to show for correlations and p-values, respectively. col.low, col.mid, col.high The color for the low (-1), mid(0) and high (1) points in the color key. Defaults to blue, white, and red, respectively. lab.x.position, lab.y.position The position of the x and y axis label. Defaults to "bottom" and "right" if

type = "lower" or "top" and "left" if type = "upper".

legend.position

The legend position in the plot.

legend.title The title of the color key. Defaults to "Pearson's Correlation".

size.text.cor The size of the text for correlation values. Defaults to 3.

size.text.signif

The size of the text for significance values (stars or p-values). Defaults to 3.

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```
size.text.lab The size of the text for labels. Defaults to 10.
... Currently not used.
```

Value

An object of class gg, ggplot

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

plot.cvalidation

Plot the RMSPD of a cross-validation procedure

Description

Boxplot showing the Root Means Square Prediction Difference of of a cross validation procedure.

Usage

```
## S3 method for class 'cvalidation'
plot(
    x,
    violin = FALSE,
    export = FALSE,
    order_box = FALSE,
    x.lab = NULL,
    y.lab = NULL,
    size.tex.lab = 12,
    file.type = "pdf",
    file.name = NULL,
    plot_theme = theme_metan(),
    width = 6,
    height = 6,
    resolution = 300,
```

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```
col.violin = "gray90",
col.boxplot = "gray70",
col.boxplot.win = "cyan",
width.boxplot = 0.6,
x.lim = NULL,
x.breaks = waiver(),
...
)
```

Arguments

X	An object of class cvalidation fitted with the functions cv_ammi(), cv_ammif(), cv_blup(), or a bound object fitted with bind_cv().	
violin	Define if a violin plot is used with boxplot. Default is 'TRUE'	
export	Export (or not) the plot. Default is T.	
order_box	Logical argument. If TRUE then the boxplots will be ordered according to the values of the RMSPD.	
x.lab	The label of x-axis. New arguments can be inserted as $x.lab = 'my \times label'$.	
y.lab	The label of y-axis. New arguments can be inserted as y.lab = 'my y label'.	
size.tex.lab	The size of the text in axis text and labels.	
file.type	The type of file to be exported. Default is pdf, Graphic can also be exported in *.tiff format by declaring file.type = 'tiff'.	
file.name	The name of the file for exportation, default is NULL, i.e. the files are automatically named.	
plot_theme	The graphical theme of the plot. Default is plot_theme = theme_metan(). For more details,see ggplot2::theme().	
width	The width 'inch' of the plot. Default is 6.	
height	The height 'inch' of the plot. Default is 6.	
resolution	The resolution of the plot. Parameter valid if file.type = 'tiff' is used. Default is 300 (300 dpi)	
col.violin	Parameter valid if violin = T. Define the color of the violin plot. Default is 'gray90.	
col.boxplot	Define the color for boxplot. Default is 'gray70'.	
col.boxplot.win		
	Define the color for boxplot of the best model. Default is 'cyan'.	
width.boxplot	The width of boxplots. Default is 0.2.	
x.lim	The range of x-axis. Default is NULL (maximum and minimum values of the data set). New arguments can be inserted as $x.lim = c(x.min, x.max)$.	
x.breaks	The breaks to be plotted in the x-axis. Default is authomatic breaks. New arguments can be inserted as x . breaks = $c(breaks)$	
	Currently not used.	

Details

Five statistics are shown in this type of plot. The lower and upper hinges correspond to the first and third quartiles (the 25th and 75th percentiles). The upper whisker extends from the hinge to the largest value no further than 1.5 * IQR from the hinge (where IQR is the inter-quartile range). The lower whisker extends from the hinge to the smallest value at most 1.5 * IQR of the hinge. Data beyond the end of the whiskers are considered outlying points.

144 plot.env_dissimilarity

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
plot.env_dissimilarity
```

Plot an object of class env_dissimilarity

Description

Create dendrograms to show the dissimilarity between environments.

Usage

```
## S3 method for class 'env_dissimilarity'
plot(x, var = 1, nclust = NULL, ...)
```

Arguments

x An object of class env_dissimilarity

var The variable to plot. Defaults to var = 1 the first variable of x.

nclust The number of clusters to show.

... Other arguments bo be passed to the function stats::hclust().

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
mod <- env_dissimilarity(data_ge, ENV, GEN, REP, GY)
plot(mod)</pre>
```

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```
plot.env_stratification
```

Plot the env_stratification model

Description

This function plots the correlation between environments generated with env_stratification()

Usage

```
## S3 method for class 'env_stratification'
plot(x, var = 1, ...)
```

Arguments

```
x An object of class env_stratification
var The variable to plot. Defaults to var = 1 the first variable of x.
... Further arguments passed to plot.corr_coef()
```

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

See Also

```
env_dissimilarity()
```

plot.fai_blup

plot.fai_blup

Multi-trait selection index

Description

Plot the multitrait index based on factor analysis and ideotype-design proposed by Rocha et al. (2018).

Usage

```
## S3 method for class 'fai_blup'
plot(
    x,
    ideotype = 1,
    SI = 15,
    radar = TRUE,
    arrange.label = FALSE,
    size.point = 2.5,
    size.line = 0.7,
    size.text = 10,
    col.sel = "red",
    col.nonsel = "black",
    ...
)
```

Arguments

X	An object of class waasb
ideotype	The ideotype to be plotted. Default is 1.
SI	An integer (0-100). The selection intensity in percentage of the total number of genotypes.
radar	Logical argument. If true (default) a radar plot is generated after using coord_polar().
arrange.label	Logical argument. If TRUE, the labels are arranged to avoid text overlapping. This becomes useful when the number of genotypes is large, say, more than 30.
size.point	The size of the point in graphic. Defaults to 2.5.
size.line	The size of the line in graphic. Defaults to 0.7.
size.text	The size for the text in the plot. Defaults to 10.
col.sel	The colour for selected genotypes. Defaults to "red".
col.nonsel	The colour for nonselected genotypes. Defaults to "black".
	Other arguments to be passed from ggplot2::theme().

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

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References

Rocha, J.R.A.S.C.R, J.C. Machado, and P.C.S. Carneiro. 2018. Multitrait index based on factor analysis and ideotype-design: proposal and application on elephant grass breeding for bioenergy. GCB Bioenergy 10:52-60. doi: 10.1111/gcbb.12443

Examples

plot.gafem

Several types of residual plots

Description

Residual plots for a output model of class gafem. Seven types of plots are produced: (1) Residuals vs fitted, (2) normal Q-Q plot for the residuals, (3) scale-location plot (standardized residuals vs Fitted Values), (4) standardized residuals vs Factor-levels, (5) Histogram of raw residuals and (6) standardized residuals vs observation order, and (7) 1:1 line plot.

Usage

```
## S3 method for class 'gafem' plot(x, ...)
```

Arguments

x An object of class gafem.

... Additional arguments passed on to the function residual_plots()

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

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Examples

```
library(metan)
model <- gafem(data_g, GEN, REP, PH)

plot(model)
plot(model,
    which = c(3, 5),
    nrow = 2,
    labels = TRUE,
    size.lab.out = 4)</pre>
```

plot.gamem

Several types of residual plots

Description

Residual plots for a output model of class gamem. Six types of plots are produced: (1) Residuals vs fitted, (2) normal Q-Q plot for the residuals, (3) scale-location plot (standardized residuals vs Fitted Values), (4) standardized residuals vs Factor-levels, (5) Histogram of raw residuals and (6) standardized residuals vs observation order. For a waasb object, normal Q-Q plot for random effects may also be obtained declaring type = 're'

Usage

```
## S3 method for class 'gamem'
plot(
  х,
  var = 1,
  type = "res",
  position = "fill",
  rotate = FALSE,
  conf = 0.95,
  out = "print",
  n.dodge = 1,
  check.overlap = FALSE,
  labels = FALSE,
  plot_theme = theme_metan(),
  alpha = 0.2,
  fill.hist = "gray",
  col.hist = "black",
  col.point = "black",
  col.line = "red",
  col.lab.out = "red",
  size.line = 0.7,
  size.text = 10,
  width.bar = 0.75,
  size.lab.out = 2.5,
  size.tex.lab = 10,
  size.shape = 1.5,
```

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```
bins = 30,
which = c(1:4),
ncol = NULL,
nrow = NULL,
...
)
```

Arguments

ncol, nrow

An object of class gamem. Х The variable to plot. Defaults to var = 1 the first variable of x. var One of the "res" to plot the model residuals (default), type = 're' to plot nortype mal Q-Q plots for the random effects, or "vcomp" to create a bar plot with the variance components. The position adjustment when type = "vcomp". Defaults to "fill", which position shows relative proportions at each trait by stacking the bars and then standardizing each bar to have the same height. Use position = "stack" to plot the phenotypic variance for each trait. Logical argument. If rotate = TRUE the plot is rotated, i.e., traits in y axis and rotate value in the x axis. Level of confidence interval to use in the Q-Q plot (0.95 by default). conf out How the output is returned. Must be one of the 'print' (default) or 'return'. n.dodge The number of rows that should be used to render the x labels. This is useful for displaying labels that would otherwise overlap. Silently remove overlapping labels, (recursively) prioritizing the first, last, and check.overlap middle labels. labels Logical argument. If TRUE labels the points outside confidence interval limits. plot_theme The graphical theme of the plot. Default is plot_theme = theme_metan(). For more details, see ggplot2::theme(). The transparency of confidence band in the Q-Q plot. Must be a number between alpha 0 (opaque) and 1 (full transparency). fill.hist The color to fill the histogram. Default is 'gray'. col.hist The color of the border of the histogram. Default is 'black'. col.point The color of the points in the graphic. Default is 'black'. col.line The color of the lines in the graphic. Default is 'red'. col.lab.out The color of the labels for the 'outlying' points. The size of the line in graphic. Defaults to 0.7. size.line size.text The size for the text in the plot. Defaults to 10. width.bar The width of the bars if type = "contribution". size.lab.out The size of the labels for the 'outlying' points. size.tex.lab The size of the text in axis text and labels. size.shape The size of the shape in the plots. bins The number of bins to use in the histogram. Default is 30. which Which graphics should be plotted. Default is which = c(1:4) that means that the first four graphics will be plotted.

The number of columns and rows of the plot pannel. Defaults to NULL

Additional arguments passed on to the function patchwork::wrap_plots().

plot.ge_effects

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

plot.ge_cluster

Plot an object of class ge_cluster

Description

Plot an object of class ge_cluster

Usage

```
## S3 method for class 'ge_cluster'
plot(x, nclust = NULL, xlab = "", ...)
```

Arguments

x An object of class ge_clusternclust The number of clusters to show.

xlab The label of the x axis.

... Other arguments passed from the function plot.hclust.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

plot.ge_effects

Plot an object of class ge_effects

Description

Plot the regression model generated by the function ge_effects.

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Usage

```
## S3 method for class 'ge_effects'
plot(
    x,
    var = 1,
    plot_theme = theme_metan(),
    x.lab = NULL,
    y.lab = NULL,
    leg.position = "right",
    size.text = 12,
    ...
)
```

Arguments

X	An object of class ge_effects
var	The variable to plot. Defaults to $var = 1$ the first variable of x.
plot_theme	The graphical theme of the plot. Default is plot_theme = theme_metan(). For more details, see ggplot2::theme().
x.lab	The label of x-axis. Each plot has a default value. New arguments can be inserted as $x.lab = "my label"$.
y.lab	The label of y-axis. Each plot has a default value. New arguments can be inserted as $y.lab = "my label"$.
leg.position	The position of the legend.
size.text	The size of the text in the axes text and labels. Default is 12.
	Current not used.

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

See Also

```
ge_plot()
```

```
library(metan)
ge_eff <- ge_effects(data_ge2, ENV, GEN, PH)
plot(ge_eff)</pre>
```

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plot.ge_factanal *F*

Plot the ge_factanal model

Description

This function plot the scores for genotypes obtained in the factor analysis to interpret the stability

Usage

```
## S3 method for class 'ge_factanal'
plot(
  Х,
  var = 1,
  plot_theme = theme_metan(),
  x.lim = NULL,
  x.breaks = waiver(),
  x.lab = NULL,
  y.lim = NULL,
  y.breaks = waiver(),
  y.lab = NULL,
  shape = 21,
  col.shape = "gray30",
  col.alpha = 1,
  size.shape = 2.2,
  size.bor.tick = 0.3,
  size.tex.lab = 12,
  size.tex.pa = 3.5,
  force.repel = 1,
  line.type = "dashed",
  line.alpha = 1,
  col.line = "black",
  size.line = 0.5,
)
```

Arguments

X	An object of class ge_factanal
var	The variable to plot. Defaults to $var = 1$ the first variable of x.
plot_theme	The graphical theme of the plot. Default is plot_theme = theme_metan(). For more details, see ggplot2::theme().
x.lim	The range of x-axis. Default is NULL (maximum and minimum values of the data set). New arguments can be inserted as $x.lim = c(x.min,x.max)$.
x.breaks	The breaks to be plotted in the x-axis. Default is authomatic breaks. New arguments can be inserted as x . breaks = $c(breaks)$
x.lab	The label of x-axis. Each plot has a default value. New arguments can be inserted as $x.lab = "my label"$.
y.lim	The range of x-axis. Default is NULL. The same arguments than $x.lim$ can be used.

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y.breaks	The breaks to be plotted in the x-axis. Default is authomatic breaks. The same arguments than x.breaks can be used.
y.lab	The label of y-axis. Each plot has a default value. New arguments can be inserted as y.lab = "my label".
shape	The shape for genotype indication in the plot. Default is 1 (circle). Values between 21–25: 21 (circle), 22 (square), 23 (diamond), 24 (up triangle), and 25 (low triangle) allows a color for fill the shape.
col.shape	The shape color for genotypes. Must be one value or a vector of colors with the same length of the number of genotypes. Default is "gray30". Other values can be attributed. For example, transparent_color(), will make a plot with only an outline around the shape area.
col.alpha	The alpha value for the color. Default is 1. Values must be between 0 (full transparency) to 1 (full color).
size.shape	The size of the shape (both for genotypes and environments). Default is 2.2.
size.bor.tick	The size of tick of shape. Default is 0.3. The size of the shape will be size.shape + size.bor.tick
size.tex.lab	The size of the text in the axes text and labels. Default is 12.
size.tex.pa	The size of the text of the plot area. Default is 3.5.
force.repel	Force of repulsion between overlapping text labels. Defaults to 1.
line.type	The type of the line that indicate the means in the biplot. Default is "solid". Other values that can be attributed are: "blank", no lines in the biplot, "dashed", "dotted", "dotdash", "longdash", and "twodash".
line.alpha	The alpha value that combine the line with the background to create the appearance of partial or full transparency. Default is 0.4. Values must be between "0" (full transparency) to "1" (full color).
col.line	The color of the line that indicate the means in the biplot. Default is "gray"
size.line	The size of the line that indicate the means in the biplot. Default is 0.5 .
	Currently not used

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

See Also

```
ge_factanal()
```

plot.ge_reg

```
plot(model)
plot(model,
    size.shape = 3,
    force.repel = 10,
    col.shape = "orange",
    col.line = "red")
```

plot.ge_reg

Plot an object of class ge_reg

Description

Plot the regression model generated by the function ge_reg.

Usage

```
## S3 method for class 'ge_reg'
plot(
    x,
    var = 1,
    type = 1,
    plot_theme = theme_metan(),
    x.lim = NULL,
    x.breaks = waiver(),
    x.lab = NULL,
    y.lim = NULL,
    y.breaks = waiver(),
    y.lab = NULL,
    leg.position = "right",
    size.tex.lab = 12,
    ...
)
```

Arguments

X	An object of class ge_factanal
var	The variable to plot. Defaults to $var = 1$ the first variable of x.
type	The type of plot to show. type = 1 produces a plot with the environmental index in the x axis and the genotype mean yield in the y axis. type = 2 produces a plot with the response variable in the x axis and the slope/deviations of the regression in the y axis.
plot_theme	The graphical theme of the plot. Default is plot_theme = theme_metan(). For more details, see ggplot2::theme().
x.lim	The range of x-axis. Default is NULL (maximum and minimum values of the data set). New arguments can be inserted as $x.lim = c(x.min,x.max)$.
x.breaks	The breaks to be plotted in the x-axis. Default is authomatic breaks. New arguments can be inserted as $x.breaks = c(breaks)$

plot.gge

x.lab	The label of x-axis. Each plot has a default value. New arguments can be inserted as $x.lab = "my label"$.
y.lim	The range of x-axis. Default is NULL. The same arguments than $x.lim$ can be used.
y.breaks	The breaks to be plotted in the x-axis. Default is authomatic breaks. The same arguments than x.breaks can be used.
y.lab	The label of y-axis. Each plot has a default value. New arguments can be inserted as y.lab = "my label".
leg.position	The position of the legend.
size.tex.lab	The size of the text in the axes text and labels. Default is 12.
	Currently not used

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

See Also

```
ge_factanal()
```

Examples

```
library(metan)
model <- ge_reg(data_ge2, ENV, GEN, REP, PH)
plot(model)</pre>
```

plot.gge

Create GGE, GT or GYT biplots

Description

Produces a ggplot2-based GGE-GT-GYT biplot based on a model fitted with the functions gge(), gtb(), and gytb().

Usage

```
## S3 method for class 'gge'
plot(
    x,
    var = 1,
    type = 1,
    sel_env = NA,
    sel_gen = NA,
    sel_gen1 = NA,
```

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```
sel_gen2 = NA,
  shape.gen = 21,
  shape.env = 23,
  line.type.gen = "dotted",
  size.shape = 2.2,
  size.shape.win = 3.2,
  size.stroke = 0.3,
  col.stroke = "black",
  col.gen = "blue",
  col.env = "forestgreen",
  col.line = "forestgreen",
  col.alpha = 1,
  col.circle = "gray",
  col.alpha.circle = 0.5,
  leg.lab = NULL,
  size.text.gen = 4,
  size.text.env = 4,
  size.text.lab = 12,
  size.text.win = 4.5,
  size.line = 0.5,
 axis_expand = 1.2,
  title = TRUE,
 plot_theme = theme_metan(),
)
```

Arguments

Х

An object with classes gge gtb, or gytb.

var

The variable to plot (useful for gge objects. Defaults to var = 1 the first variable of x.

type

The type of biplot to produce.

- 1. Basic biplot.
- 2. Mean performance vs. stability (gge biplots) or the The Average Tester Coordination view for genotype-trait and genotype-yield*trait biplots.
- 3. Which-won-where.
- 4. Discriminativeness vs. representativeness.
- 5. Examine an environment (or trait/yield*trait combination).
- 6. Ranking environments (or trait/yield*trait combination).
- 7. Examine a genotype.
- 8. Ranking genotypes.
- 9. Compare two genotypes.
- 10. Relationship among environments (or trait/yield*trait combination).

sel_env, sel_gen

The name of the environment (or trait/yield*trait combination) and genotype to examine when type = 5 and type = 7, respectively. Must be a string which matches a environment or genotype label.

```
sel_gen1, sel_gen2
```

The name of genotypes to compare between when type = 9. Must be a string present in the genotype's name.

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shape.gen, shape.env

The shape for genotype and environment indication in the biplot. Defaults to shape.gen = 21 (circle) for genotypes and shape.env = 23 (rhombus) for environments. Values must be between 21-25: 21 (circle), 22 (square), 23 (rhombus), 24 (up triangle), and 25 (low triangle).

line.type.gen The line type to highlith the genotype's vectors. Defaults to line.type.gen == "dotted."

size. shape The size of the shape (both for genotypes and environments). Defaults to 2.2.

size.shape.win The size of the shape for winners genotypes when type = 3. Defaults to 3.2. size.stroke, col.stroke

The width and color of the border, respectively. Default to size.stroke = 0.3 and col.stroke = "black". The size of the shape will be size.shape + size.stroke

col.gen, col.env, col.line

Color for genotype/environment labels and for the line that passes through the biplot origin. Defaults to col.gen = 'blue', col.env = 'forestgreen', and col.line = 'forestgreen'.

col.alpha The alpha value for the color. Defaults to 1. Values must be between 0 (full transparency) to 1 (full color).

col.circle, col.alpha.circle

The color and alpha values for the circle lines. Defaults to 'gray' and 0.4, respectively.

leg.lab The labs of legend. Defaults to NULL is c('Env', 'Gen').

size.text.gen, size.text.env, size.text.lab

The size of the text for genotypes, environments and labels, respectively.

size.text.win The text size to use for winner genotypes where type = 3 and for the two selected genotypes where type = 9. Defaults to 4.5.

size.line The size of the line in biplots (Both for segments and circles).

axis_expand multiplication factor to expand the axis limits by to enable fitting of labels. De-

faults to 1.2

title Logical values (Defaults to TRUE) to include automatically generated informa-

tion in the plot such as singular value partitioning, scaling and centering.

plot_theme The graphical theme of the plot. Default is plot_theme = theme_metan(). For

more details, see ggplot2::theme().

... Currently not used.

Value

A ggplot2-based biplot.

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Yan, W., and M.S. Kang. 2003. GGE biplot analysis: a graphical tool for breeders, geneticists, and agronomists. CRC Press.

158 plot.mgidi

Examples

```
library(metan)
mod <- gge(data_ge, ENV, GEN, GY)
plot(mod)
plot(mod,
    type = 2,
    col.gen = 'blue',
    col.env = 'red',
    size.text.gen = 2)</pre>
```

plot.mgidi

Plot the multi-trait genotype-ideotype distance index

Description

Makes a radar plot showing the multi-trait genotype-ideotype distance index

Usage

```
## S3 method for class 'mgidi'
plot(
  х,
  SI = 15,
  radar = TRUE,
  type = "index",
  position = "fill",
  rotate = FALSE,
  genotypes = "selected",
  n.dodge = 1,
  check.overlap = FALSE,
  x.lab = NULL,
  y.lab = NULL,
  title = NULL,
  arrange.label = FALSE,
  size.point = 2.5,
  size.line = 0.7,
  size.text = 10,
  width.bar = 0.75,
  col.sel = "red",
  col.nonsel = "gray",
  legend.position = "bottom",
)
```

Arguments

x An object of class mgidi
SI An integer (0-100). The selection intensity in percentage of the total number of genotypes.

radar Logical argument. If true (default) a radar plot is generated after using coord_polar().

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type	The type of the plot. Defaults to "index". Use type = "contribution" to show the contribution of each factor to the MGIDI index of the selected genotypes/treatments.
position	The position adjustment when type = "contribution". Defaults to "fill", which shows relative proportions at each trait by stacking the bars and then standardizing each bar to have the same height. Use position = "stack" to plot the MGIDI index for each genotype/treatment.
rotate	Logical argument. If rotate = TRUE the plot is rotated, i.e., traits in y axis and value in the x axis.
genotypes	When type = "contribution" defines the genotypes to be shown in the plot. By default (genotypes = "selected" only selected genotypes are shown. Use genotypes = "all" to plot the contribution for all genotypes.)
n.dodge	The number of rows that should be used to render the x labels. This is useful for displaying labels that would otherwise overlap.
check.overlap	Silently remove overlapping labels, (recursively) prioritizing the first, last, and middle labels.
x.lab, y.lab	The labels for the axes x and y, respectively. x label is set to null when a radar plot is produced.
title	The plot title when type = "contribution".
arrange.label	Logical argument. If TRUE, the labels are arranged to avoid text overlapping. This becomes useful when the number of genotypes is large, say, more than 30.
size.point	The size of the point in graphic. Defaults to 2.5.
size.line	The size of the line in graphic. Defaults to 0.7.
size.text	The size for the text in the plot. Defaults to 10.
width.bar	The width of the bars if type = "contribution". Defaults to 0.75.
col.sel	The colour for selected genotypes. Defaults to "red".
col.nonsel	The colour for nonselected genotypes. Defaults to "gray".
legend.position	
	The position of the legend.
	Other arguments to be passed from ggplot2::theme().

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

160 plot.mtmps

plot.mtmps

Plot the multi-trait stability index

Description

Makes a radar plot showing the multitrait stability index proposed by Olivoto et al. (2019)

Usage

```
## S3 method for class 'mtmps'
plot(
  Х,
  SI = 15,
  type = "index",
  position = "fill",
  genotypes = "selected",
  title = TRUE,
  radar = TRUE,
  arrange.label = FALSE,
  x.lab = NULL,
  y.lab = NULL,
  size.point = 2.5,
  size.line = 0.7,
  size.text = 10,
  width.bar = 0.75,
  n.dodge = 1,
  check.overlap = FALSE,
  invert = FALSE,
  col.sel = "red",
  col.nonsel = "black",
  legend.position = "bottom",
)
```

Arguments

x	An object computed with mps().
SI	An integer (0-100). The selection intensity in percentage of the total number of genotypes.
type	The type of the plot. Defaults to "index". Use type = "contribution" to show the contribution of each factor to the MTMPS index of the selected genotypes.
position	The position adjustment when type = "contribution". Defaults to "fill", which shows relative proportions at each trait by stacking the bars and then standardizing each bar to have the same height. Use position = "stack" to plot the MGIDI index for each genotype.
genotypes	When type = "contribution" defines the genotypes to be shown in the plot. By default (genotypes = "selected" only selected genotypes are shown. Use genotypes = "all" to plot the contribution for all genotypes.)
title	Logical values (Defaults to TRUE) to include automatically generated titles.

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radar	Logical argument. If true (default) a radar plot is generated after using coord_polar().
arrange.label	Logical argument. If TRUE, the labels are arranged to avoid text overlapping. This becomes useful when the number of genotypes is large, say, more than 30.
x.lab, y.lab	The labels for the axes x and y, respectively. x label is set to null when a radar plot is produced.
size.point	The size of the point in graphic. Defaults to 2.5.
size.line	The size of the line in graphic. Defaults to 0.7.
size.text	The size for the text in the plot. Defaults to 10.
width.bar	The width of the bars if type = "contribution". Defaults to 0.75.
n.dodge	The number of rows that should be used to render the x labels. This is useful for displaying labels that would otherwise overlap.
check.overlap	Silently remove overlapping labels, (recursively) prioritizing the first, last, and middle labels.
invert	Logical argument. If TRUE, rotate the plot.
col.sel	The colour for selected genotypes. Defaults to "red".
col.nonsel	The colour for nonselected genotypes. Defaults to "black".
legend.position	
	The position of the legend.
	Other arguments to be passed from ggplot2::theme().

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Olivoto, T., A.D.C. L\'ucio, J.A.G. da silva, B.G. Sari, and M.I. Diel. 2019. Mean performance and stability in multi-environment trials II: Selection based on multiple traits. Agron. J. (in press).

```
library(metan)
model <-
mps(data_ge,
    env = ENV,
    gen = GEN,
    rep = REP,
    resp = everything())
selection <- mtmps(model)
plot(selection)</pre>
```

162 plot.mtsi

plot.mtsi

Plot the multi-trait stability index

Description

Makes a radar plot showing the multitrait stability index proposed by Olivoto et al. (2019)

Usage

```
## S3 method for class 'mtsi'
plot(
  Х,
  SI = 15,
  type = "index",
  position = "fill",
  genotypes = "selected",
  title = TRUE,
  radar = TRUE,
  arrange.label = FALSE,
  x.lab = NULL,
  y.lab = NULL,
  size.point = 2.5,
  size.line = 0.7,
  size.text = 10,
  width.bar = 0.75,
  n.dodge = 1,
  check.overlap = FALSE,
  invert = FALSE,
  col.sel = "red",
  col.nonsel = "black",
  legend.position = "bottom",
)
```

Arguments

x	An object of class mtsi
SI	An integer (0-100). The selection intensity in percentage of the total number of genotypes.
type	The type of the plot. Defaults to "index". Use type = "contribution" to show the contribution of each factor to the MGIDI index of the selected genotypes.
position	The position adjustment when type = "contribution". Defaults to "fill", which shows relative proportions at each trait by stacking the bars and then standardizing each bar to have the same height. Use position = "stack" to plot the MGIDI index for each genotype.
genotypes	When type = "contribution" defines the genotypes to be shown in the plot. By default (genotypes = "selected" only selected genotypes are shown. Use genotypes = "all" to plot the contribution for all genotypes.)
title	Logical values (Defaults to TRUE) to include automatically generated titles.

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radar	Logical argument. If true (default) a radar plot is generated after using coord_polar().
arrange.label	Logical argument. If TRUE, the labels are arranged to avoid text overlapping. This becomes useful when the number of genotypes is large, say, more than 30.
x.lab, y.lab	The labels for the axes x and y, respectively. x label is set to null when a radar plot is produced.
size.point	The size of the point in graphic. Defaults to 2.5.
size.line	The size of the line in graphic. Defaults to 0.7.
size.text	The size for the text in the plot. Defaults to 10.
width.bar	The width of the bars if type = "contribution". Defaults to 0.75.
n.dodge	The number of rows that should be used to render the x labels. This is useful for displaying labels that would otherwise overlap.
check.overlap	Silently remove overlapping labels, (recursively) prioritizing the first, last, and middle labels.
invert	Logical argument. If TRUE, rotate the plot.
col.sel	The colour for selected genotypes. Defaults to "red".
col.nonsel	The colour for nonselected genotypes. Defaults to "black".
legend.position	
	The position of the legend.
	Other arguments to be passed from ggplot2::theme().

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Olivoto, T., A.D.C. L\'ucio, J.A.G. da silva, B.G. Sari, and M.I. Diel. 2019. Mean performance and stability in multi-environment trials II: Selection based on multiple traits. Agron. J. (in press).

```
library(metan)
mtsi_model <- waasb(data_ge, ENV, GEN, REP, resp = c(GY, HM))
mtsi_index <- mtsi(mtsi_model)
plot(mtsi_index)</pre>
```

plot.resp_surf

```
plot.performs_ammi
```

Several types of residual plots

Description

Residual plots for a output model of class performs_ammi. Seven types of plots are produced: (1) Residuals vs fitted, (2) normal Q-Q plot for the residuals, (3) scale-location plot (standardized residuals vs Fitted Values), (4) standardized residuals vs Factor-levels, (5) Histogram of raw residuals and (6) standardized residuals vs observation order, and (7) 1:1 line plot.

Usage

```
## S3 method for class 'performs_ammi'
plot(x, ...)
```

Arguments

x An object of class performs_ammi.

... Additional arguments passed on to the function residual_plots()

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
model <- performs_ammi(data_ge, ENV, GEN, REP, GY)
plot(model)
plot(model,
    which = c(3, 5),
    nrow = 2,
    labels = TRUE,
    size.lab.out = 4)</pre>
```

plot.resp_surf

Plot the response surface model

Description

Plot the response surface model using a contour plot

plot.resp_surf

Usage

```
## S3 method for class 'resp_surf'
plot(
    x,
    xlab = NULL,
    ylab = NULL,
    resolution = 100,
    bins = 10,
    plot_theme = theme_metan(),
    ...
)
```

Arguments

X	An object of class resp_surf
xlab, ylab	The label for the x and y axis, respectively. Defaults to original variable names.
resolution	The resolution of the contour plot. Defaults to 100. higher values produce high-resolution plots but may increase the computation time.
bins	The number of bins shown in the plot. Defaults to 10.
plot_theme	The graphical theme of the plot. Default is plot_theme = theme_metan(). For more details, see ggplot2::theme().
	Currently not used

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

plot.sh

plot.sh Plot the Smith-Hazel index

Description

Makes a radar plot showing the individual genetic worth for the Smith-Hazel index

Usage

```
## S3 method for class 'sh'
plot(
    x,
    SI = 15,
    radar = TRUE,
    arrange.label = FALSE,
    size.point = 2.5,
    size.line = 0.7,
    size.text = 10,
    col.sel = "red",
    col.nonsel = "black",
    ...
)
```

Arguments

X	An object of class sh
SI	An integer (0-100). The selection intensity in percentage of the total number of genotypes.
radar	Logical argument. If true (default) a radar plot is generated after using coord_polar().
arrange.label	Logical argument. If TRUE, the labels are arranged to avoid text overlapping. This becomes useful when the number of genotypes is large, say, more than 30.
size.point	The size of the point in graphic. Defaults to 2.5.
size.line	The size of the line in graphic. Defaults to 0.7.
size.text	The size for the text in the plot. Defaults to 10.
col.sel	The colour for selected genotypes. Defaults to "red".
col.nonsel	The colour for nonselected genotypes. Defaults to "black".
	Other arguments to be passed from ggplot2::theme().

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

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Examples

```
library(metan)
vcov <- covcor_design(data_g, GEN, REP, everything())
means <- as.matrix(vcov$means)
pcov <- vcov$phen_cov
gcov <- vcov$geno_cov

index <- Smith_Hazel(means, pcov = pcov, gcov = gcov, weights = rep(1, 15))
plot(index)</pre>
```

plot.waas

Several types of residual plots

Description

Residual plots for a output model of class waas. Seven types of plots are produced: (1) Residuals vs fitted, (2) normal Q-Q plot for the residuals, (3) scale-location plot (standardized residuals vs Fitted Values), (4) standardized residuals vs Factor-levels, (5) Histogram of raw residuals and (6) standardized residuals vs observation order, and (7) 1:1 line plot.

Usage

```
## S3 method for class 'waas' plot(x, ...)
```

Arguments

x An object of class waas.

... Additional arguments passed on to the function residual_plots()

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

```
library(metan)
model <- waas(data_ge, ENV, GEN, REP, GY)
plot(model)
plot(model,
    which = c(3, 5),
    nrow = 2,
    labels = TRUE,
    size.lab.out = 4)</pre>
```

168 plot.waasb

plot.waasb

Several types of residual plots

Description

Residual plots for a output model of class waas and waasb. Six types of plots are produced: (1) Residuals vs fitted, (2) normal Q-Q plot for the residuals, (3) scale-location plot (standardized residuals vs Fitted Values), (4) standardized residuals vs Factor-levels, (5) Histogram of raw residuals and (6) standardized residuals vs observation order. For a waasb object, normal Q-Q plot for random effects may also be obtained declaring type = 're'

Usage

```
## S3 method for class 'waasb'
plot(
  Х,
  var = 1,
  type = "res",
  position = "fill",
  rotate = FALSE,
  conf = 0.95,
  out = "print",
  n.dodge = 1,
  check.overlap = FALSE,
  labels = FALSE,
  plot_theme = theme_metan(),
  alpha = 0.2,
  fill.hist = "gray",
  col.hist = "black"
  col.point = "black",
  col.line = "red",
  col.lab.out = "red",
  size.line = 0.7,
  size.text = 10,
  width.bar = 0.75,
  size.lab.out = 2.5,
  size.tex.lab = 10,
  size.shape = 1.5,
  bins = 30,
  which = c(1:4),
  ncol = NULL,
  nrow = NULL,
)
```

Arguments

x An object of class waasb.

var The variable to plot. Defaults to var = 1 the first variable of x.

plot.waasb 169

type One of the "res" to plot the model residuals (default), type = 're' to plot normal Q-Q plots for the random effects, or "vcomp" to create a bar plot with the variance components. position The position adjustment when type = "vcomp". Defaults to "fill", which shows relative proportions at each trait by stacking the bars and then standardizing each bar to have the same height. Use position = "stack" to plot the phenotypic variance for each trait. rotate Logical argument. If rotate = TRUE the plot is rotated, i.e., traits in y axis and value in the x axis. Level of confidence interval to use in the Q-Q plot (0.95 by default). conf How the output is returned. Must be one of the 'print' (default) or 'return'. out The number of rows that should be used to render the x labels. This is useful for n.dodge displaying labels that would otherwise overlap. check.overlap Silently remove overlapping labels, (recursively) prioritizing the first, last, and middle labels. labels Logical argument. If TRUE labels the points outside confidence interval limits. plot_theme The graphical theme of the plot. Default is plot_theme = theme_metan(). For more details, see ggplot2::theme(). alpha The transparency of confidence band in the Q-Q plot. Must be a number between 0 (opaque) and 1 (full transparency). fill.hist The color to fill the histogram. Default is 'gray'. The color of the border of the histogram. Default is 'black'. col.hist col.point The color of the points in the graphic. Default is 'black'. col.line The color of the lines in the graphic. Default is 'red'. col.lab.out The color of the labels for the 'outlying' points. size.line The size of the line in graphic. Defaults to 0.7. The size for the text in the plot. Defaults to 10. size.text width.bar The width of the bars if type = "contribution". size.lab.out The size of the labels for the 'outlying' points. The size of the text in axis text and labels. size.tex.lab size.shape The size of the shape in the plots. bins The number of bins to use in the histogram. Default is 30. which Which graphics should be plotted. Default is which = c(1:4) that means that the first four graphics will be plotted. The number of columns and rows of the plot pannel. Defaults to NULL ncol, nrow Additional arguments passed on to the function patchwork::wrap_plots().

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

170 plot.wsmp

Examples

plot.wsmp

Plot heat maps with genotype ranking

Description

Plot heat maps with genotype ranking in two ways.

Usage

```
## S3 method for class 'wsmp'
plot(x, var = 1, type = 1, y.lab = NULL, x.lab = NULL, size.lab = 12, ...)
```

Arguments

x	The object returned by the function wsmp.
var	The variable to plot. Defaults to $var = 1$ the first variable of x.
type	1 = Heat map Ranks: this graphic shows the genotype ranking considering the WAASB index estimated with different numbers of Principal Components; 2 = Heat map WAASY-GY ratio: this graphic shows the genotype ranking considering the different combinations in the WAASB/GY ratio.
y.lab	The label of y axis. Default is 'Genotypes'.
x.lab	The label of x axis. Default is 'Number of axes'.
size.lab	The size of the
	Currently not used.

Details

The first type of heatmap shows the genotype ranking depending on the number of principal component axis used for estimating the WAASB index. The second type of heatmap shows the genotype ranking depending on the WAASB/GY ratio. The ranks obtained with a ratio of 100/0 considers exclusively the stability for the genotype ranking. On the other hand, a ratio of 0/100 considers exclusively the productivity for the genotype ranking. Four clusters of genotypes are shown by label colors (red) unproductive and unstable genotypes; (blue) productive, but unstable genotypes; (black) stable, but unproductive genotypes; and (green), productive and stable genotypes.

Value

An object of class gg.

plot_blup 171

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

plot_blup

Plot the BLUPs for genotypes

Description

[Stable]

Plot the predicted BLUP of the genotypes.

Usage

```
plot_blup(
  х,
  var = 1,
  which = "gen",
  ncol = NULL,
  nrow = NULL,
  prob = 0.05,
  export = FALSE,
  file.type = "pdf",
  file.name = NULL,
  plot_theme = theme_metan(),
  width = 6,
  height = 6,
  err.bar = TRUE,
  size.err.bar = 0.5,
  size.shape = 3.5,
  size.tex.lab = 12,
  height.err.bar = 0.3,
  x.lim = NULL,
  x.breaks = waiver(),
  col.shape = c("blue", "red"),
  y.lab = "Genotypes",
```

172 plot_blup

```
x.lab = NULL,
n.dodge = 1,
check.overlap = FALSE,
panel.spacing = 0.15,
resolution = 300,
...
)
```

Arguments

Ę	guinents	
	x	The waasb object
	var	The variable to plot. Defaults to $var = 1$ the first variable of x.
	which	Which plot to shown. If which = "gen" (default) plots the BLUPs for genotypes. To create a plot showing the BLUPs for genotype-environment combinations, use which = "ge".
	ncol, nrow	The number of columns and rows, respectively, to be shown in the plot when $which = "ge"$.
	prob	The probability error for constructing confidence interval.
	export	Export (or not) the plot. Default is TRUE.
	file.type	If export = TRUE, define the type of file to be exported. Default is pdf, Graphic can also be exported in *.tiff format by declaring file.type = "tiff".
	file.name	The name of the file for exportation, default is NULL, i.e. the files are automatically named.
	plot_theme	The graphical theme of the plot. Default is plot_theme = theme_metan(). For more details, see ggplot2::theme().
	width	The width "inch" of the plot. Default is 6.
	height	The height "inch" of the plot. Default is 6.
	err.bar	Logical value to indicate if an error bar is shown. Defaults to TRUE.
	size.err.bar	The size of the error bar for the plot. Default is 0.5.
	size.shape	The size of the shape (both for genotypes and environments). Default is 3.5.
	size.tex.lab	The size of the text in axis text and labels.
	height.err.bar	The height for error bar. Default is 0.3.
	x.lim	The range of x-axis. Default is NULL (maximum and minimum values of the data set). New arguments can be inserted as $x.lim = c(x.min,x.max)$.
	x.breaks	The breaks to be plotted in the x-axis. Default is authomatic breaks. New arguments can be inserted as $x.breaks = c(breaks)$
	col.shape	A vector of length 2 that contains the color of shapes for genotypes above and below of the mean, respectively. Default is c("blue", "red").
	y.lab	The label of the y-axis in the plot. Default is "Genotypes".
	x.lab	The label of the x-axis in the plot. Default is $NULL$, i.e., the name of the selected variable.
	n.dodge	The number of rows that should be used to render the Y labels. This is useful for displaying labels that would otherwise overlap.
	check.overlap	Silently remove overlapping labels, (recursively) prioritizing the first, last, and middle labels.

panel.spacing Defines the spacing between panels when which = "ge".

plot_ci 173

```
resolution The resolution of the plot. Parameter valid if file.type = "tiff" is used. Default is 300 (300 dpi)
... Currently not used.
```

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

See Also

```
plot_scores(), plot_waasby()
```

Examples

plot_ci

Plot the confidence interval for correlation

Description

[Stable]

This function plots the 95% confidence interval for Pearson's correlation coefficient generated by the function corr_ci.

Usage

```
plot_ci(
  object,
  fill = NULL,
  position.fill = 0.3,
  x.lab = NULL,
  y.lab = NULL,
  y.lim = NULL,
  y.breaks = waiver(),
  shape = 21,
  col.shape = "black",
```

plot_ci

```
fill.shape = "orange",
  size.shape = 2.5,
  width.errbar = 0.2,
  main = TRUE,
  invert.axis = TRUE,
  reorder = TRUE,
  legend.position = "bottom",
  plot_theme = theme_metan()
)
```

Arguments

object	An object generate by the function corr_ci()
fill	If corr_ci() is computed with the argument by use fill to fill the shape by each level of the grouping variable by.
position.fill	The position of shapes and errorbar when fill is used. Defaults to 0.3 .
x.lab	The label of x-axis, set to 'Pairwise combinations'. New arguments can be inserted as $x.lab = \text{'my label'}$.
y.lab	The label of y-axis, set to 'Pearson's correlation coefficient' New arguments can be inserted as y.lab = 'my label'.
y.lim	The range of x-axis. Default is NULL. The same arguments than ${\tt x.lim}$ can be used.
y.breaks	The breaks to be plotted in the x-axis. Default is authomatic breaks. The same arguments than x.breaks can be used.
shape	The shape point to represent the correlation coefficient. Default is 21 (circle). Values must be between 21–25: 21 (circle), 22 (square), 23 (diamond), 24 (up triangle), and 25 (low triangle).
col.shape	The color for the shape edge. Set to black.
fill.shape	The color to fill the shape. Set to orange.
size.shape	The size for the shape point. Set to 2.5.
width.errbar	The width for the errorbar showing the CI.
main	The title of the plot. Set to main = FALSE to ommite the plot title.
invert.axis	Should the names of the pairwise correlation appear in the y-axis?
reorder	Logical argument. If TRUE (default) the pairwise combinations are reordered according to the correlation coefficient.
legend.positio	
	The position of the legend when using fill argument. Defaults to "bottom".
plot_theme	The graphical theme of the plot. Default is plot_theme = theme_metan(). For more details, see ggplot2::theme().

Value

An object of class gg, ggplot.

plot_eigen 175

Examples

plot_eigen

Plot the eigenvalues

Description

[Stable]

Plot the eigenvalues for from singular value decomposition of BLUP interaction effects matrix.

Usage

```
plot_eigen(
  х,
  var = 1,
  export = FALSE,
  plot_theme = theme_metan(),
  file.type = "pdf",
  file.name = NULL,
  width = 6,
  height = 6,
  size.shape = 3.5,
  size.line = 1,
  size.tex.lab = 12,
  y.lab = "Eigenvalue",
  y2.lab = "Accumulated variance",
  x.lab = "Number of multiplicative terms",
  resolution = 300,
)
```

plot_eigen

Arguments

x	The waasb object
var	The variable to plot. Defaults to $var = 1$ the first variable of x.
export	Export (or not) the plot. Default is TRUE.
plot_theme	The graphical theme of the plot. Default is plot_theme = theme_metan(). For more details, see ggplot2::theme().
file.type	If export = TRUE, define the type of file to be exported. Default is pdf, Graphic can also be exported in *.tiff format by declaring file.type = "tiff".
file.name	The name of the file for exportation, default is NULL, i.e. the files are automatically named.
width	The width "inch" of the plot. Default is 6.
height	The height "inch" of the plot. Default is 6.
size.shape	The size of the shape. Default is 3.5.
size.line	The size of the line. Default is 1.
size.tex.lab	The size of the text in axis text and labels.
y.lab	The label of the y-axis in the plot. Default is "Eigenvalue".
y2.lab	The label of the second y-axis in the plot. Default is "Accumulated variance".
x.lab	The label of the x-axis in the plot. Default is "Number of multiplicative terms".
resolution	The resolution of the plot. Parameter valid if file.type = "tiff" is used. Default is $300 (300 \text{ dpi})$
	Currently not used.

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

See Also

```
plot_scores(), plot_waasby()
```

plot_scores

Plot scores in different graphical interpretations

Description

[Stable]

Plot scores of genotypes and environments in different graphical interpretations.

Biplots type 1 and 2 are well known in AMMI analysis. In the plot type 3, the scores of both genotypes and environments are plotted considering the response variable and the WAASB, an stability index that considers all significant principal component axis of traditional AMMI models or all principal component axis estimated with BLUP-interaction effects (Olivoto et al. 2019). Plot type 4 may be used to better understand the well known 'which-won-where' pattern, facilitating the recommendation of appropriate genotypes targeted for specific environments, thus allowing the exploitation of narrow adaptations.

Usage

```
plot_scores(
  Х,
  var = 1,
  type = 1,
  first = "PC1",
  second = "PC2",
  repel = TRUE,
  repulsion = 1,
  polygon = FALSE,
  title = TRUE,
  plot_theme = theme_metan(),
  axis.expand = 1.1,
  x.lim = NULL,
  y.lim = NULL,
  x.breaks = waiver(),
  y.breaks = waiver(),
  x.lab = NULL,
  y.lab = NULL,
  shape.gen = 21,
  shape.env = 23,
  size.shape.gen = 2.2,
  size.shape.env = 2.2,
  size.shape = deprecated(),
  size.bor.tick = 0.3,
  size.tex.lab = 12,
  size.tex.pa = deprecated(),
  size.tex.gen = 3.5,
  size.tex.env = 3.5,
  size.line = 0.5,
  size.segm.line = 0.5,
  col.bor.gen = "black",
  col.bor.env = "black",
  col.line = "black",
```

```
col.gen = "blue",
  col.env = "forestgreen",
  col.alpha.gen = NA,
  col.alpha.env = NA,
  col.segm.gen = transparent_color(),
  col.segm.env = "forestgreen",
 highlight = NULL,
  col.highlight = "red",
  leg.lab = c("Env", "Gen"),
  line.type = "solid",
  line.alpha = 0.9,
  resolution = 300,
  file.type = "pdf",
  export = FALSE,
  file.name = NULL,
 width = 8,
 height = 7,
 color = TRUE,
)
```

Arguments

x An object fitted with the functions performs_ammi(), waas(), waas_means(), or waasb().

var The variable to plot. Defaults to var = 1 the first variable of x.

type type of biplot to produce

• type = 1 The default. Produces an AMMI1 biplot (Y x PC1) to make inferences related to stability and productivity.

- type = 2 Produces an AMMI2 biplot (PC1 x PC2) to make inferences related to the interaction effects. Use the arguments first or second to change the default IPCA shown in the plot.
- type = 3 Valid for objects of class waas or waasb, produces a biplot showing the GY x WAASB.
- type = 4 Produces a plot with the Nominal yield x Environment PC.

first, second The IPCA to be shown in the first (x) and second (y) axis. By default, IPCA1 is

shown in the x axis and IPCA2 in the y axis. For example, use second = "PC3"

to shown the IPCA3 in the y axis.

repel If TRUE (default), the text labels repel away from each other and away from the

data points.

repulsion Force of repulsion between overlapping text labels. Defaults to 1.

polygon Logical argument. If TRUE, a polygon is drawn when type = 2.

title Logical values (Defaults to TRUE) to include automatically generated titles

plot_theme The graphical theme of the plot. Default is plot_theme = theme_metan(). For

more details, see ggplot2::theme().

axis.expand Multiplication factor to expand the axis limits by to enable fitting of labels.

Default is 1.1.

x.lim, y.lim The range of x and y axes, respectively. Default is NULL (maximum and mini-

mum values of the data set). New values can be inserted as x.lim = c(x.min, x.max)

or y.lim = c(y.min, y.max).

x.breaks, y.breaks

The breaks to be plotted in the x and y axes, respectively. Defaults to waiver() (automatic breaks). New values can be inserted, for example, as x.breaks = c(0.1, 0.2, 0.3) or x.breaks = seq(0.1, by = 0.2)

x.lab, y.lab The label of x and y axes, respectively. Defaults to NULL, i.e., each plot has a default axis label. New values can be inserted as x.lab = 'my label'.

shape.gen, shape.env

The shape for genotypes and environments indication in the biplot. Default is 21 (circle) for genotypes and 23 (diamond) for environments. Values must be between 21–25: 21 (circle), 22 (square), 23 (diamond), 24 (up triangle), and 25 (low triangle).

size.shape.gen, size.shape.env

The size of the shapes for genotypes and environments respectively. Defaults to 2.2.

size.shape [Deprecated]

size.bor.tick The size of tick of shape. Default is 0.3. The size of the shape will be max(size.shape.gen, size.sh + size.bor.tick

size.tex.lab, size.tex.gen, size.tex.env

The size of the text for axis labels (Defaults to 12), genotypes labels, and environments labels (Defaults to 3.5).

size.tex.pa [Deprecated]

size.line The size of the line that indicate the means in the biplot. Default is 0.5.

size.segm.line The size of the segment that start in the origin of the biplot and end in the scores values. Default is 0.5.

col.bor.gen, col.bor.env

The color of the shape's border for genotypes and environments, respectively.

col.line The color of the line that indicate the means in the biplot. Default is 'gray'

col.gen, col.env

The shape color for genotypes (Defaults to 'blue') and environments ('forestgreen'). Must be length one or a vector of colors with the same length of the number of genotypes/environments.

col.alpha.gen, col.alpha.env

The alpha value for the color for genotypes and environments, respectively. Defaults to NA. Values must be between 0 (full transparency) to 1 (full color).

col.segm.gen, col.segm.env

The color of segment for genotypes (Defaults to transparent_color()) and environments (Defaults to 'forestgreen'), respectively. Valid arguments for plots with type = 1 or type = 2 graphics.

highlight Genotypes/environments to be highlight in the plot. Defaults to NULL.

col.highlight The color for shape/labels when a value is provided in highlight.

leg. lab The labs of legend. Default is Gen and Env.

line.type The type of the line that indicate the means in the biplot. Default is 'solid'.

Other values that can be attributed are: 'blank', no lines in the biplot, 'dashed', 'dotted', 'dotdash', 'longdash', and 'twodash'.

line.alpha The alpha value that combine the line with the background to create the appearance of partial or full transparency. Default is 0.4. Values must be between '0' (full transparency) to '1' (full color).

resoluti	The resolution of the plot. Parameter valid if file.type = 'tiff' is used. Default is 300 (300 dpi)
file.typ	The type of file to be exported. Valid parameter if export = T TRUE. Default is 'pdf'. The graphic can also be exported in *.tiff format by declaring file.type = 'tiff'.
export	Export (or not) the plot. Default is FALSE.
file.nam	The name of the file for exportation, default is NULL, i.e. the files are automatically named.
width	The width 'inch' of the plot. Default is 8.
height	The height 'inch' of the plot. Default is 7.
color	Should type 4 plot have colors? Default to TRUE.
	Currently not used.

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Olivoto, T., A.D.C. L\'ucio, J.A.G. da silva, V.S. Marchioro, V.Q. de Souza, and E. Jost. 2019. Mean performance and stability in multi-environment trials I: Combining features of AMMI and BLUP techniques. Agron. J. 111:2949-2960. doi: 10.2134/agronj2019.03.0220

See Also

```
plot_eigen()
```

```
library(metan)
# AMMI model
model <- waas(data_ge,</pre>
            env = ENV,
            gen = GEN,
            rep = REP,
            resp = everything())
# GY x PC1 for variable GY (default plot)
plot_scores(model)
# PC1 x PC2 (variable HM)
plot_scores(model,
           polygon = TRUE, # Draw a convex hull polygon
           var = "HM",
                                      # or var = 2 to select variable
           highlight = c("G1", "G2"), # Highlight genotypes 2 and 3
           type = 2)
                                      # type of biplot
# PC3 x PC4 (variable HM)
```

plot_waasby 181

```
# Change size of plot fonts and colors
# Minimal theme
plot_scores(model,
           var = "HM",
           type = 2,
           first = "PC3"
           second = "PC4",
           col.gen = "black",
           col.env = "gray",
           col.segm.env = "gray",
           size.tex.gen = 5,
           size.tex.env = 2,
           size.tex.lab = 16,
           plot_theme = theme_metan_minimal())
# WAASB index
waasb_model <- waasb(data_ge, ENV, GEN, REP, GY)</pre>
# GY x WAASB
# Highlight genotypes 2 and 8
plot_scores(waasb_model,
            type = 3,
            highlight = c("G2", "G8"))
```

plot_waasby

Plot WAASBY values for genotype ranking

Description

[Stable]

Plot heat maps with genotype ranking in two ways.

```
plot_waasby(
  х,
  var = 1,
  export = F,
  file.type = "pdf",
  file.name = NULL,
  plot_theme = theme_metan(),
  width = 6,
  height = 6,
  size.shape = 3.5,
  size.tex.lab = 12,
  col.shape = c("blue", "red"),
  x.lab = "WAASBY",
  y.lab = "Genotypes",
  x.breaks = waiver(),
  resolution = 300,
)
```

182 plot_waasby

Arguments

х	The WAASBY object
var	The variable to plot. Defaults to $var = 1$ the first variable of x.
export	Export (or not) the plot. Default is T.
file.type	The type of file to be exported. Default is pdf, Graphic can also be exported in *.tiff format by declaring file.type = "tiff".
file.name	The name of the file for exportation, default is NULL, i.e. the files are automatically named.
plot_theme	The graphical theme of the plot. Default is plot_theme = theme_metan(). For more details, see ggplot2::theme().
width	The width "inch" of the plot. Default is 8.
height	The height "inch" of the plot. Default is 7.
size.shape	The size of the shape in the plot. Default is 3.5.
size.tex.lab	The size of the text in axis text and labels.
col.shape	A vector of length 2 that contains the color of shapes for genotypes above and below of the mean, respectively. Default is c("blue", "red").
x.lab	The label of the x axis in the plot. Default is "WAASBY".
y.lab	The label of the y axis in the plot. Default is "Genotypes".
x.breaks	The breaks to be plotted in the x-axis. Default is authomatic breaks. New arguments can be inserted as x . breaks = $c(breaks)$
resolution	The resolution of the plot. Parameter valid if file.type = "tiff" is used. Default is 300 (300 dpi)
	Currently not used.

Value

An object of class gg, ggplot.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

See Also

```
plot_scores()
```

predict.gamem 183

predict.gamem

Predict method for gamem fits

Description

Obtains predictions from an object fitted with gamem().

Usage

```
## S3 method for class 'gamem'
predict(object, ...)
```

Arguments

object An object of class gamem
... Currently not used

Value

A tibble with the predicted values for each variable in the model

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

184 predict.gge

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nrea	1C	Т	000

Predict a two-way table based on GGE model

Description

Predict the means for a genotype-vs-environment trial based on a Genotype plus Genotype-vs-Environment interaction (GGE) model.

Usage

```
## S3 method for class 'gge'
predict(object, naxis = 2, output = "wide", ...)
```

Arguments

object An object of class gge.

naxis The the number of principal components to be used in the prediction. Generally,

two axis may be used. In this case, the estimated values will be those shown in

the biplot.

output The type of output. It must be one of the 'long' (default) returning a long-

format table with the columns for environment (ENV), genotypes (GEN) and response variable (Y); or 'wide' to return a two-way table with genotypes in

the row, environments in the columns, filled by the estimated values.

... Currently not used.

Details

This function is used to predict the response variable of a two-way table (for examples the yielding of g genotypes in e environments) based on GGE model. This prediction is based on the number of principal components used. For more details see Yan and Kang (2007).

Value

A two-way table with genotypes in rows and environments in columns if output = "wide" or a long format (columns ENV, GEN and Y) if output = "long" with the predicted values by the GGE model.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Yan, W., and M.S. Kang. 2003. GGE biplot analysis: a graphical tool for breeders, geneticists, and agronomists. CRC Press.

predict.performs_ammi 185

Examples

```
library(metan)
mod <- gge(data_ge, GEN, ENV, c(GY, HM))
predict(mod)</pre>
```

predict.performs_ammi Predict the means of a performs_ammi object

Description

Predict the means of a performs_ammi object considering a specific number of axis.

Usage

```
## S3 method for class 'performs_ammi'
predict(object, naxis = 2, ...)
```

Arguments

object An object of class performs_ammi

naxis The the number of axis to be use in the prediction. If object has more than one

variable, then naxis must be a vector.

... Additional parameter for the function

Details

This function is used to predict the response variable of a two-way table (for examples the yielding of the i-th genotype in the j-th environment) based on AMMI model. This prediction is based on the number of multiplicative terms used. If naxis = 0, only the main effects (AMMI0) are used. In this case, the predicted mean will be the predicted value from OLS estimation. If naxis = 1 the AMMI1 (with one multiplicative term) is used for predicting the response variable. If naxis = min(gen-1;env-1), the AMMIF is fitted and the predicted value will be the cell mean, i.e. the mean of R-replicates of the i-th genotype in the j-th environment. The number of axis to be used must be carefully chosen. Procedures based on Postdictive success (such as Gollobs's d.f.) or Predictive success (such as cross-validation) should be used to do this. This package provide both. performs_ammi() function compute traditional AMMI analysis showing the number of significant axis. On the other hand, cv_ammif() function provide a cross-validation, estimating the RMSPD of all AMMI-family models, based on resampling procedures.

Value

A list where each element is the predicted values by the AMMI model for each variable.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

186 predict.waas

Examples

predict.waas

Predict the means of a waas object

Description

Predict the means of a waas object considering a specific number of axis.

Usage

```
## S3 method for class 'waas'
predict(object, naxis = 2, ...)
```

Arguments

object An object of class waas

naxis The the number of axis to be use in the prediction. If object has more than one

variable, then naxis must be a vector.

... Additional parameter for the function

Details

This function is used to predict the response variable of a two-way table (for examples the yielding of the i-th genotype in the j-th environment) based on AMMI model. This prediction is based on the number of multiplicative terms used. If naxis = 0, only the main effects (AMMI0) are used. In this case, the predicted mean will be the predicted value from OLS estimation. If naxis = 1 the AMMI1 (with one multiplicative term) is used for predicting the response variable. If naxis = min(gen-1;env-1), the AMMIF is fitted and the predicted value will be the cell mean, i.e. the mean of R-replicates of the i-th genotype in the j-th environment. The number of axis to be used must be carefully chosen. Procedures based on Postdictive success (such as Gollobs's d.f.) or Predictive success (such as cross-validation) should be used to do this. This package provide both. waas() function compute traditional AMMI analysis showing the number of significant axis. On the other hand, $cv_ammif()$ function provide a cross-validation, estimating the RMSPD of all AMMI-family models, based on resampling procedures.

Value

A list where each element is the predicted values by the AMMI model for each variable.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

predict.waasb 187

Examples

predict.waasb

Predict method for waasb fits

Description

Obtains predictions from an object fitted with waasb().

Usage

```
## S3 method for class 'waasb'
predict(object, ...)
```

Arguments

object An object of class waasb
... Currently not used

Value

A tibble with the predicted values for each variable in the model

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

188 print.Annicchiarico

print.AMMI_indexes
Print an object of class AMMI_indexes

Description

Print the AMMI_indexes object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory into a *.txt file.

Usage

```
## S3 method for class 'AMMI_indexes'
print(x, which = "stats", export = FALSE, file.name = NULL, digits = 3, ...)
```

Arguments

x	An object of class AMMI_indexes.	
which	Which should be printed. Defaults to "stats". Other possible values are "ranks" for genotype ranking and "ssi" for the simultaneous selection index.	
export	A logical argument. If TRUE, a *.txt file is exported to the working directory.	
file.name	The name of the file if export = TRUE	
digits	The significant digits to be shown.	
	Options used by the tibble package to format the output. See tibble::print() for more details.	

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

print. Annicchiarico Print an object of class Annicchiarico

Description

Print the Annicchiarico object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory into a *.txt file.

```
## S3 method for class 'Annicchiarico'
print(x, export = FALSE, file.name = NULL, digits = 3, ...)
```

print.anova_ind 189

Arguments

X	The Annicchiarico x
export	A logical argument. If TRUE, a *.txt file is exported to the working directory.
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
• • •	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
Ann <- Annicchiarico(data_ge2,
  env = ENV,
  gen = GEN,
  rep = REP,
  resp = PH
)
print(Ann)</pre>
```

print.anova_ind

Print an object of class anova_ind

Description

Print the anova_ind object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory into a *.txt file.

Usage

```
## S3 method for class 'anova_ind'
print(x, export = FALSE, file.name = NULL, digits = 3, ...)
```

Arguments

X	An object of class anova_ind.
export	A logical argument. If TRUE, a *.txt file is exported to the working directory.
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

190 print.anova_joint

Examples

```
library(metan)
model <- data_ge %>% anova_ind(ENV, GEN, REP, c(GY, HM))
print(model)
```

print.anova_joint

Print an object of class anova_joint

Description

Print the anova_joint object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory into a *.txt file.

Usage

```
## S3 method for class 'anova_joint'
print(x, export = FALSE, file.name = NULL, digits = 3, ...)
```

Arguments

x	An object of class anova_joint.
export	A logical argument. If TRUE, a *.txt file is exported to the working directory.
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

```
library(metan)
model <- data_ge %>% anova_joint(ENV, GEN, REP, c(GY, HM))
print(model)
```

print.can_cor 191

nrı	nt	can	cor

Print an object of class can_cor

Description

Print an object of class can_cor object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory.

Usage

```
## S3 method for class 'can_cor'
print(x, export = FALSE, file.name = NULL, digits = 3, ...)
```

Arguments

X	An object of class can_cor.
export	A logical argument. If TRUE T, a *.txt file is exported to the working directory
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
•••	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

print.coincidence

Print an object of class coincidence

Description

Print a coincidence object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory.

```
## S3 method for class 'coincidence'
print(x, export = FALSE, file.name = NULL, digits = 4, ...)
```

192 print.colindiag

Arguments

X	An object of class coincidence.	
export	A logical argument. If TRUE, a *.txt file is exported to the working directory	
file.name	The name of the file if export = TRUE	
digits	The significant digits to be shown.	
• • •	Options used by the tibble package to format the output. See tibble::print() for more details.	

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
sel1 <- paste("G", 1:30, sep = "")
sel2 <- paste("G", 16:45, sep = "")
coinc <- coincidence_index(sel1 = sel1, sel2 = sel2, total = 150)
print(coinc)</pre>
```

print.colindiag

Print an object of class colindiag

Description

Print the colindiag object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory into a *.txt file.

Usage

```
## S3 method for class 'colindiag'
print(x, export = FALSE, file.name = NULL, digits = 3, ...)
```

Arguments

X	The object of class colindiag
export	A logical argument. If TRUE, a *.txt file is exported to the working directory.
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
•••	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

print.corr_coef 193

Examples

```
library(metan)
col <- colindiag(data_ge2)
print(col)</pre>
```

print.corr_coef

Print an object of class corr_coef

Description

Print the corr_coef object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory into a *.txt file.

Usage

```
## S3 method for class 'corr_coef'
print(x, export = FALSE, file.name = NULL, digits = 3, ...)
```

Arguments

X	An object of class corr_coef
export	A logical argument. If TRUE, a *.txt file is exported to the working directory.
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
	Options used by the tibble package to format the output. See tibble::formatting() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

```
library(metan)
sel <- corr_coef(data_ge2, EP, EL, CD, CL)
print(sel)</pre>
```

	-
print.	ecovalence

Print an object of class ecovalence

Description

Print the ecovalence object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory into a *.txt file.

Usage

```
## S3 method for class 'ecovalence'
print(x, export = FALSE, file.name = NULL, digits = 3, ...)
```

Arguments

X	The ecovalence x
export	A logical argument. If TRUE, a *.txt file is exported to the working directory.
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
•••	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
print.env_dissimilarity
```

Print an object of class env_dissimilarity

Description

Print the env_dissimilarity object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory into a *.txt file.

```
## S3 method for class 'env_dissimilarity'
print(x, export = FALSE, file.name = NULL, digits = 3, ...)
```

print.env_stratification 195

Arguments

x An object of class env_dissimilarity.

export A logical argument. If TRUE, a *.txt file is exported to the working directory.

file.name The name of the file if export = TRUE digits The significant digits to be shown.

... Currently not used.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
mod <- env_dissimilarity(data_ge, ENV, GEN, REP, GY)
print(mod)</pre>
```

```
print.env_stratification
```

Print the env_stratification model

Description

Print an object of class ge_factanal in two ways. By default, the results are shown in the R console. The results can also be exported to the directory.

Usage

```
## S3 method for class 'env_stratification'
print(x, export = FALSE, file.name = NULL, digits = 3, ...)
```

Arguments

x An object of class env_stratification.

export A logical argument. If TRUE, a *.txt file is exported to the working directory

file.name The name of the file if export = TRUE digits The significant digits to be shown.

... Currently not used.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

196 print.Fox

Examples

print.Fox

Print an object of class Fox

Description

Print the Fox object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory into a *.txt file.

Usage

```
## S3 method for class 'Fox'
print(x, export = FALSE, file.name = NULL, digits = 3, ...)
```

Arguments

x	The Fox x
export	A logical argument. If TRUE, a *.txt file is exported to the working directory.
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

```
library(metan)
library(metan)
out <- Fox(data_ge2, ENV, GEN, PH)
print(out)</pre>
```

print.gamem 197

print.gamem	Print an object of class gamem

Description

Print the gamem object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory.

Usage

```
## S3 method for class 'gamem'
print(x, export = FALSE, file.name = NULL, digits = 4, ...)
```

Arguments

X	An object fitted with the function gamem().
export	A logical argument. If TRUE, a *.txt file is exported to the working directory
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
alpha <- gamem(data_alpha,
  gen = GEN,
  rep = REP,
  block = BLOCK,
  resp = YIELD
)
print(alpha)</pre>
```

print.ge_factanal

Print an object of class ge_factanal

Description

Print the ge_factanal object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory.

198 print.ge_reg

Usage

```
## S3 method for class 'ge_factanal'
print(x, export = FALSE, file.name = NULL, digits = 4, ...)
```

Arguments

X	An object of class ge_factanal.
export	A logical argument. If TRUE, a *.txt file is exported to the working directory
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
model <- ge_factanal(data_ge2,
  env = ENV,
  gen = GEN,
  rep = REP,
  resp = PH
)
print(model)</pre>
```

print.ge_reg

Print an object of class ge_reg

Description

Print the ge_reg object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory into a *.txt file.

Usage

```
## S3 method for class 'ge_reg'
print(x, export = FALSE, file.name = NULL, digits = 3, ...)
```

Arguments

X	An object of class ge_reg.
export	A logical argument. If TRUE, a *.txt file is exported to the working directory.
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
	Options used by the tibble package to format the output. See tibble::print() for more details.

print.ge_stats

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Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
model <- ge_reg(data_ge2, ENV, GEN, REP, PH)
print(model)</pre>
```

print.ge_stats

Print an object of class ge_stats

Description

Print the ge_stats object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory into a *.txt file.

Usage

```
## S3 method for class 'ge_stats'
print(x, what = "all", export = FALSE, file.name = NULL, digits = 3, ...)
```

Arguments

	A 12 4 6 1
Х	An object of class ge_stats.
what	What should be printed. what = "all" for both statistics and ranks, what = "stats" for statistics, and what = "ranks" for ranks.
export	A logical argument. If TRUE, a *.txt file is exported to the working directory.
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

```
library(metan)
model <- ge_stats(data_ge, ENV, GEN, REP, GY)
print(model)</pre>
```

200 print.lpcor

print.Huehn	Print an object ofclass Huehn
p. 2	1

Description

Print the Huehn object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory into a *.txt file.

Usage

```
## S3 method for class 'Huehn'
print(x, export = FALSE, file.name = NULL, digits = 3, ...)
```

Arguments

X	An object of class Huehn.
export	A logical argument. If TRUE, a *.txt file is exported to the working directory.
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
model <- Huehn(data_ge2, ENV, GEN, PH)
print(model)</pre>
```

print.lpcor

Print the partial correlation coefficients

Description

Print an object of class 1pcor or or 1pcor_group in two ways. By default, the results are shown in the R console. The results can also be exported to the directory.

```
## S3 method for class 'lpcor'
print(x, export = FALSE, file.name = NULL, digits = 3, ...)
```

print.mgidi 201

Arguments

x	An object of class 1pcor or 1pcor_group.
export	A logical argument. If TRUE, a *.txt file is exported to the working directory
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

print.mgidi Print an object of class mgidi Print a mgidi object in two ways. By default, the results are shown in the R console. The results can also be

exported to the directory.

Description

Print an object of class mgidi Print a mgidi object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory.

Usage

```
## S3 method for class 'mgidi'
print(x, export = FALSE, file.name = NULL, digits = 4, ...)
```

Arguments

X	An object of class mgidi.
export	A logical argument. If TRUE T, a *.txt file is exported to the working directory
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
	Options used by the tibble package to format the output. See tibble::print() for more details.

202 print.mtmps

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

print.mtmps

Print an object of class mtmps

Description

Print a mtmps object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory.

Usage

```
## S3 method for class 'mtmps'
print(x, export = FALSE, file.name = NULL, digits = 4, ...)
```

Arguments

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

```
library(metan)
model <-
mps(data_ge,
    env = ENV,
    gen = GEN,
    rep = REP,
    resp = everything())
selection <- mtmps(model)
print(selection)</pre>
```

print.mtsi 203

print.mtsi	Print an object of class mtsi	

Description

Print a mtsi object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory.

Usage

```
## S3 method for class 'mtsi'
print(x, export = FALSE, file.name = NULL, digits = 4, ...)
```

Arguments

X	An object of class mtsi.
export	A logical argument. If TRUE T, a *.txt file is exported to the working directory
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

```
library(metan)
# Based on stability only
MTSI_MODEL <- waasb(data_ge,
  resp = c(GY, HM),
  gen = GEN,
  env = ENV,
  rep = REP
)
MTSI_index <- mtsi(MTSI_MODEL)
print(MTSI_index)</pre>
```

print.path_coeff

print.path_coeff	Print an object of class path_coeff

Description

Print an object generated by the function 'path_coeff()'. By default, the results are shown in the R console. The results can also be exported to the directory.

Usage

```
## S3 method for class 'path_coeff'
print(x, export = FALSE, file.name = NULL, digits = 4, ...)
```

Arguments

X	An object of class path_coeff or group_path.
export	A logical argument. If TRUE, a *.txt file is exported to the working directory
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

print.performs_ammi 205

Description

Print the performs_ammi object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory.

Usage

```
## S3 method for class 'performs_ammi'
print(x, export = FALSE, file.name = NULL, digits = 4, ...)
```

Arguments

X	An object of class performs_ammi.
export	A logical argument. If TRUE, a *.txt file is exported to the working directory
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

print.Schmildt

Print an object of class Schmildt

Description

Print the Schmildt object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory into a *.txt file.

```
## S3 method for class 'Schmildt'
print(x, export = FALSE, file.name = NULL, digits = 3, ...)
```

206 print.sh

Arguments

Х	The Schmildt x
export	A logical argument. If TRUE, a *.txt file is exported to the working directory.
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
	Options used by the tibble package to format the output. See tibble::formatting() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

print.sh

Print an object of class sh

Description

Print a sh object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory.

Usage

```
## S3 method for class 'sh'
print(x, export = FALSE, file.name = NULL, digits = 4, ...)
```

Arguments

X	An object of class sh.
export	A logical argument. If TRUE, a *.txt file is exported to the working directory
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

print.Shukla 207

Examples

```
vcov <- covcor_design(data_g, GEN, REP, everything())
means <- as.matrix(vcov$means)
pcov <- vcov$phen_cov
gcov <- vcov$geno_cov

index <- Smith_Hazel(means, pcov = pcov, gcov = gcov, weights = rep(1, 15))
print(index)</pre>
```

print.Shukla

Print an object of class Shukla

Description

Print the Shukla object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory into a *.txt file.

Usage

```
## S3 method for class 'Shukla'
print(x, export = FALSE, file.name = NULL, digits = 3, ...)
```

Arguments

```
x The Shukla x
export A logical argument. If TRUE, a *.txt file is exported to the working directory.
file.name The name of the file if export = TRUE
digits The significant digits to be shown.
... Options used by the tibble package to format the output. See tibble::print() for more details.
```

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

```
library(metan)
eco <- Shukla(data_ge2,
   env = ENV,
   gen = GEN,
   rep = REP,
   resp = PH
)
print(eco)</pre>
```

208 print.Thennarasu

print.superiority	Print an object of class superiority	
-------------------	--------------------------------------	--

Description

Print the superiority object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory into a *.txt file.

Usage

```
## S3 method for class 'superiority'
print(x, export = FALSE, file.name = NULL, digits = 3, ...)
```

Arguments

X	An object of class superiority.
export	A logical argument. If TRUE, a *.txt file is exported to the working directory.
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
•••	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
model <- superiority(data_ge2, ENV, GEN, PH)
print(model)</pre>
```

```
print.Thennarasu
```

Print an object of class Thennarasu

Description

Print the Thennarasu object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory into a *.txt file.

```
## S3 method for class 'Thennarasu'
print(x, export = FALSE, file.name = NULL, digits = 3, ...)
```

print.waas 209

Arguments

x	An object of class Thennarasu.
export	A logical argument. If TRUE, a *.txt file is exported to the working directory.
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
	Options used by the tibble package to format the output. See tibble::print() for more details

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
model <- Thennarasu(data_ge2, ENV, GEN, PH)
print(model)</pre>
```

print.waas Print an object of class waas

Description

Print the waas object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory.

Usage

```
## S3 method for class 'waas'
print(x, export = FALSE, file.name = NULL, digits = 4, ...)
```

Arguments

Х	An object of class waas.
export	A logical argument. If TRUE, a *.txt file is exported to the working directory
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
•••	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

210 print.waasb

Examples

```
library(metan)
model <- waas(data_ge,
  resp = c(GY, HM),
  gen = GEN,
  env = ENV,
  rep = REP
)
print(model)</pre>
```

print.waasb

Print an object of class waasb

Description

Print a waasb object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory.

Usage

```
## S3 method for class 'waasb'
print(x, export = FALSE, blup = FALSE, file.name = NULL, digits = 4, ...)
```

Arguments

x	An object of class waasb.
export	A logical argument. If TRUE T, a *.txt file is exported to the working directory
blup	A logical argument. If TRUE T, the blups are shown.
file.name	The name of the file if export = TRUE
digits	The significant digits to be shown.
•••	Options used by the tibble package to format the output. See tibble::print() for more details.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

```
library(metan)
model <- waasb(data_ge,
  resp = c(GY, HM),
  gen = GEN,
  env = ENV,
  rep = REP
)
print(model)</pre>
```

print.waas_means 211

print.waas_means	Print an object of class waas_means	
------------------	-------------------------------------	--

Description

Print the waas_means object in two ways. By default, the results are shown in the R console. The results can also be exported to the directory.

Usage

```
## S3 method for class 'waas_means'
print(x, export = FALSE, file.name = NULL, digits = 4, ...)
```

Arguments

```
An object of class waas_means.

export A logical argument. If TRUE, a *.txt file is exported to the working directory file.name The name of the file if export = TRUE

digits The significant digits to be shown. See tibble::print() for more details.

Currently not used.
```

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

reorder_cormat

Reorder a correlation matrix

Description

[Stable]

Reorder the correlation matrix according to the correlation coefficient by using helust for hierarchical clustering order. This is useful to identify the hidden pattern in the matrix.

```
reorder_cormat(x)
```

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Arguments

Х

The correlation matrix

Value

The ordered correlation matrix

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
cor_mat <- corr_coef(data_ge2, PH, EH, CD, CL, ED, NKR)
cor_mat$cor
reorder_cormat(cor_mat$cor)</pre>
```

resca

Rescale a variable to have specified minimum and maximum values

Description

[Stable]

Helper function that rescales a continuous variable to have specified minimum and maximum values.

The function rescale a continuous variable as follows:

```
Rv_i = (Nmax - Nmin)/(Omax - Omin) * (O_i - Omax) + Nmax
```

Where Rv_i is the rescaled value of the ith position of the variable/vector; Nmax and Nmin are the new maximum and minimum values; OmaxandOmin are the maximum and minimum values of the original data, and O_i is the ith value of the original data.

There are basically two options to use resca to rescale a variable. The first is passing a data frame to .data argument and selecting one or more variables to be scaled using The function will return the original variables in .data plus the rescaled variable(s) with the prefix _res. By using the function group_by from **dplyr** package it is possible to rescale the variable(s) within each level of the grouping factor. The second option is pass a numeric vector in the argument values. The output, of course, will be a numeric vector of rescaled values.

```
resca(
   .data = NULL,
   ...,
   values = NULL,
   new_min = 0,
   new_max = 100,
   na.rm = TRUE,
   keep = TRUE
)
```

residual_plots 213

Arguments

.data	The dataset. Grouped data is allowed.
	Comma-separated list of unquoted variable names that will be rescaled.
values	Optional vector of values to rescale
new_min	The minimum value of the new scale. Default is 0.
new_max	The maximum value of the new scale. Default is 100
na.rm	Remove NA values? Default to TRUE.
keep	Should all variables be kept after rescaling? If false, only rescaled variables will be kept.

Value

A numeric vector if values is used as input data or a tibble if a data frame is used as input in .data.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

```
library(metan)
library(dplyr)
# Rescale a numeric vector
resca(values = c(1:5))
# Using a data frame
head(
 resca(data_ge, GY, HM, new_min = 0, new_max = 1)
)
# Rescale within factors;
# Select variables that stats with 'N' and ends with 'L';
# Compute the mean of these variables by ENV and GEN;
# Rescale the variables that ends with 'L' whithin ENV;
data_ge2 %>%
  select(ENV, GEN, starts_with("N"), ends_with("L")) %>%
 means_by(ENV, GEN) %>%
 group_by(ENV) %>%
 resca(ends_with("L")) %>%
  head(n = 13)
```

214 residual_plots

Description

[Stable]

Residual plots for a output model of class performs_ammi, waas, anova_ind, and anova_joint. Seven types of plots are produced: (1) Residuals vs fitted, (2) normal Q-Q plot for the residuals, (3) scale-location plot (standardized residuals vs Fitted Values), (4) standardized residuals vs Factorlevels, (5) Histogram of raw residuals and (6) standardized residuals vs observation order, and (7) 1:1 line plot

Usage

```
residual_plots(
  х,
  var = 1,
  conf = 0.95,
  labels = FALSE,
  plot_theme = theme_metan(),
  band.alpha = 0.2,
  point.alpha = 0.8,
  fill.hist = "gray",
  col.hist = "black",
  col.point = "black",
  col.line = "red",
  col.lab.out = "red",
  size.lab.out = 2.5,
  size.tex.lab = 10,
  size.shape = 1.5,
  bins = 30,
  which = c(1:4),
 ncol = NULL,
  nrow = NULL,
)
```

Arguments

X	An object of class performs_ammi, waas, anova_joint, or gafem	
var	The variable to plot. Defaults to $var = 1$ the first variable of x.	
conf	Level of confidence interval to use in the Q-Q plot (0.95 by default).	
labels	Logical argument. If TRUE labels the points outside confidence interval limits.	
plot_theme	The graphical theme of the plot. Default is plot_theme = theme_metan(). For more details, see ggplot2::theme().	
band.alpha, point.alpha		
	The transparency of confidence band in the Q-Q plot and the points, respectively. Must be a number between 0 (opaque) and 1 (full transparency).	
fill.hist	The color to fill the histogram. Default is 'gray'.	
col.hist	The color of the border of the histogram. Default is 'black'.	
col.point	The color of the points in the graphic. Default is 'black'.	
col.line	The color of the lines in the graphic. Default is 'red'.	
col.lab.out	The color of the labels for the 'outlying' points.	

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size.lab.out The size of the labels for the 'outlying' points.

size.tex.lab The size of the text in axis text and labels.

size.shape The size of the shape in the plots.

bins The number of bins to use in the histogram. Default is 30.

which Which graphics should be plotted. Default is which = c(1:4) that means that the first four graphics will be plotted.

ncol, nrow The number of columns and rows of the plot pannel. Defaults to NULL

Additional arguments passed on to the function patchwork::wrap_plots().

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
model <- performs_ammi(data_ge, ENV, GEN, REP, GY)</pre>
# Default plot
plot(model)
# Normal Q-Q plot
# Label possible outliers
plot(model,
     which = 2,
     labels = TRUE)
# Residual vs fitted,
# Normal Q-Q plot
# Histogram of raw residuals
# All in one row
plot(model,
     which = c(1, 2, 5),
     nrow = 1)
```

resp_surf

Response surface model

Description

[Stable]

Compute a surface model and find the best combination of factor1 and factor2 to obtain the stationary point.

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Usage

```
resp_surf(
   .data,
   factor1,
   factor2,
   rep = NULL,
   resp,
   prob = 0.05,
   verbose = TRUE
)
```

Arguments

.data	The dataset containing the columns related to Environments, factor1, factor2, replication/block and response variable(s).
factor1	The first factor, for example, dose of Nitrogen.
factor2	The second factor, for example, dose of potassium.
rep	The name of the column that contains the levels of the replications/blocks, if a designed experiment was conducted. Defaults to NULL.
resp	The response variable(s).
prob	The probability error.
verbose	If verbose = TRUE then some results are shown in the console.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Schmildt 217

|--|

Description

[Stable]

Stability analysis using the known genotypic confidence index (Annicchiarico, 1992) modified by Schmildt et al. 2011.

Usage

```
Schmildt(.data, env, gen, rep, resp, prob = 0.05, verbose = TRUE)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s)
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
rep	The name of the column that contains the levels of the replications/blocks
resp	The response variable(s). To analyze multiple variables in a single procedure use, for example, $resp = c(var1, var2, var3)$.
prob	The probability of error assumed.
verbose	Logical argument. If verbose = FALSE the code will run silently.

Value

A list where each element is the result for one variable and contains the following data frames:

- **environments** Contains the mean, environmental index and classification as favorables and unfavorables environments.
- general Contains the genotypic confidence index considering all environments.
- favorable Contains the genotypic confidence index considering favorable environments.
- unfavorable Contains the genotypic confidence index considering unfavorable environments.

Author(s)

Tiago Olivoto, <tiagoolivoto@gmail.com>

References

Annicchiarico, P. 1992. Cultivar adaptation and recommendation from alfalfa trials in Northern Italy. J. Genet. Breed. 46:269-278.

Schmildt, E.R., A.L. Nascimento, C.D. Cruz, and J.A.R. Oliveira. 2011. Avaliacao de metodologias de adaptabilidade e estabilidade de cultivares milho. Acta Sci. - Agron. 33:51-58. doi: 10.4025/actasciagron.v33i1.5817

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See Also

```
superiority(), ecovalence(), ge_stats(), Annicchiarico()
```

Examples

Select_helper

Select helper

Description

These functions allow you to select variables based operations with prefixes and suffixes and length of names.

- difference_var(): Select variables that start with a prefix AND NOT end wiht a suffix.
- intersect_var(): Select variables that start with a prefix **AND** end wiht a suffix.
- union_var(): Select variables that start with a prefix **OR** end wiht a suffix.
- width_of(): Select variables with width of n.
- width_greater_than(): Select variables with width greater than n.
- width_less_than(): Select variables with width less than n.
- lower_case_only(): Select variables that contains lower case only (e.g., "env").
- upper_case_only(): Select variables that contains upper case only (e.g., "ENV").
- title_case_only(): Select variables that contains upper case in the first character only (e.g., "Env").

Usage

```
difference_var(prefix, suffix)
intersect_var(prefix, suffix)
union_var(prefix, suffix)
width_of(n, vars = peek_vars(fn = "width_of"))
width_greater_than(n, vars = peek_vars(fn = "width_greater_than"))
width_less_than(n, vars = peek_vars(fn = "width_less_than"))
lower_case_only(vars = peek_vars(fn = "lower_case_only"))
```

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```
upper_case_only(vars = peek_vars(fn = "upper_case_only"))
title_case_only(vars = peek_vars(fn = "title_case_only"))
```

Arguments

prefix A prefix that start the variable name.

Suffix A suffix that end the variable name.

The length of variable names to select. For width_of() the selected variables contains n characters. For width_greater_than() and width_less_than() the selected variables contains greater and less characteres than n, respectively.

Vars A character vector of variable names. When called from inside selecting functions like select_cols() these are automatically set to the names of the table.

```
library(metan)
```

```
# Select variables that start with "C" and not end with "D".
data_ge2 %>%
select_cols(difference_var("C", "D"))
# Select variables that start with "C" and end with "D".
data_ge2 %>%
select_cols(intersect_var("C", "D"))
# Select variables that start with "C" or end with "D".
data_ge2 %>%
select_cols(union_var("C", "D"))
# Select variables with width name of 4
data_ge2 %>%
select_cols(width_of(4))
# Select variables with width name greater than 2
data_ge2 %>%
select_cols(width_greater_than(2))
# Select variables with width name less than 3
data_ge2 %>%
select_cols(width_less_than(3))
# Creating data with messy column names
df <- head(data_ge, 3)</pre>
colnames(df) <- c("Env", "gen", "Rep", "GY", "hm")</pre>
select_cols(df, lower_case_only())
select_cols(df, upper_case_only())
select_cols(df, title_case_only())
```

220 select_pred

select_pred	Selects a best subset of predictor variables.

Description

Selects among a set of covariates the best set of npred predictors for a given response trait resp based on AIC values.

Usage

```
select_pred(.data, resp, covariates = NULL, npred)
```

Arguments

.data A data frame with the response variable and covariates.

resp The response variable.

covariates The covariates. Defaults to NULL. In this case, all numeric traits in .data,

except that in resp are selected. To select specific covariates from .data, use a list of unquoted comma-separated variable names (e.g. traits = c(var1, var2, var3)), an specific range of variables, (e.g. traits = c(var1:var3)), or even a

select helper like starts_with("N").

npred An integer specifying the size of the subset of predictors to be selected

Value

A list with the following elements:

- sel_mod An object of class 1m that is the selected model.
- predictors The name of the selected predictors.
- AIC The Akaike's Information Criterion for the selected model.
- pred_models The Akaike's Information Criterion and the predictors selected in each step.
- predicted The predicted values considering the model in sel_mod.

Author(s)

```
Tiago Olivoto <tiagoolivoto@gmail.com>
```

```
library(metan)
mod <- select_pred(data_ge2, resp = PH, npred = 10)
mod$predictors
mod$AIC</pre>
```

Shukla 221

Shukla	Shukla's stability variance parameter	

Description

[Stable]

The function computes the Shukla's stability variance parameter (1972) and uses the Kang's non-parametric stability (rank sum) to imcorporate the mean performance and stability into a single selection criteria.

Usage

```
Shukla(.data, env, gen, rep, resp, verbose = TRUE)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
rep	The name of the column that contains the levels of the replications/blocks.
resp	The response variable(s). To analyze multiple variables in a single procedure use, for example, $resp = c(var1, var2, var3)$.
verbose	Logical argument. If verbose = FALSE the code will run silently.

Value

An object of class Shukla, which is a list containing the results for each variable used in the argument resp. For each variable, a tibble with the following columns is returned.

- **GEN** the genotype's code.
- Y the mean for the response variable.
- ShuklaVar The Shukla's stability variance parameter.
- rMean The rank for Y (decreasing).
- rShukaVar The rank for ShukaVar.
- ssiShukaVar The simultaneous selection index (ssiShukaVar = rMean + rShukaVar).

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Shukla, G.K. 1972. Some statistical aspects of partitioning genotype-environmental components of variability. Heredity. 29:238-245. doi: 10.1038/hdy.1972.87

Kang, M.S., and H.N. Pham. 1991. Simultaneous Selection for High Yielding and Stable Crop Genotypes. Agron. J. 83:161. doi: 10.2134/agronj1991.00021962008300010037x

Smith_Hazel

Examples

Smith_Hazel

Smith-Hazel index

Description

[Stable]

Computes the Smith (1936) and Hazel (1943) index given economic weights and phenotypic and genotypic variance-covariance matrices. The Smith-Hazel index is computed as follows:

$$\mathbf{b} = \mathbf{P^{-1}Aw}$$

where P and G are phenotypic and genetic covariance matrices, respectively, and b and w are vectors of index coefficients and economic weightings, respectively.

The genetic worth I of an individual genotype based on traits x, y, ..., n, is calculated as:

$$I = b_x G_x + b_y G_y + \ldots + b_n G_n$$

where b the index coefficient for the traits x, y, ..., n, respectively, and G is the individual genotype BLUPs for the traits x, y, ..., n, respectively.

Usage

```
Smith_Hazel(
   .data,
   use_data = "blup",
   pcov = NULL,
   gcov = NULL,
   SI = 15,
   weights = NULL
)
```

Arguments

.data

The input data. It can be either a two-way table with genotypes in rows and traits in columns, or an object fitted with the function gamem(). Please, see **Details** for more details.

use_data

Define which data to use If .data is an object of class gamem. Defaults to "blup" (the BLUPs for genotypes). Use "pheno" to use phenotypic means instead BLUPs for computing the index.

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pcov, gcov	The phenotypic and genotypic variance-covariance matrix, respectively. De-
	faults to NULL. If a two-way table is informed in . data these matrices are manda-
	forv

SI The selection intensity (percentage). Defaults to 20

The vector of economic weights. Defaults to a vector of 1s with the same length weights

of the number of traits.

Details

When using the phenotypic means in .data, be sure the genotype's code are in rownames. If .data is an object of class gamem them the BLUPs for each genotype are used to compute the index. In this case, the genetic covariance components are estimated by mean cross products.

Value

An object of class hz containing:

- **b**: the vector of index coefficient.
- index: The genetic worth.
- sel_dif_trait: The selection differencial.
- sel_gen: The selected genotypes.
- gcov: The genotypic variance-covariance matrix
- pcov: The phenotypic variance-covariance matrix

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

```
Smith, H.F. 1936. A discriminant function for plant selection. Ann. Eugen. 7:240-250. doi: 10.1111/
j.14691809.1936.tb02143.x
```

Hazel, L.N. 1943. The genetic basis for constructing selection indexes. Genetics 28:476-90. PMID:17247099

See Also

```
mtsi(), mgidi(), fai_blup()
```

```
vcov <- covcor_design(data_g, GEN, REP, everything())</pre>
means <- as.matrix(vcov$means)</pre>
pcov <- vcov$phen_cov</pre>
gcov <- vcov$geno_cov</pre>
index <- Smith_Hazel(means, pcov = pcov, gcov = gcov, weights = rep(1, 15))</pre>
```

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solve_svd

Pseudoinverse of a square matrix

Description

[Stable]

This function computes the Moore-Penrose pseudoinverse of a square matrix using singular value decomposition.

Usage

```
solve_svd(x, tolerance = 2.220446e-16)
```

Arguments

x A square matrix

tolerance The tolerance to consider an eigenvalue equals to zero.

Value

A matrix with the same dimension of x.

Author(s)

Tiago Olivoto, <tiagoolivoto@gmail.com>

Examples

```
library(metan)
mat <- matrix(c(1, 4, 2, 8), ncol = 2)
det(mat)
solve_svd(mat)</pre>
```

split_factors

Split a data frame by factors

Description

[Stable]

Split a data frame into subsets grouping by one or more factors.

This function is used to split a data frame into a named list where each element is a level of the grouping variable (or combination of grouping variables).

Usage

```
split_factors(.data, ..., keep_factors = FALSE)
as.split_factors(.data, keep_factors = FALSE)
is.split_factors(x)
```

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Arguments

.data	The data that will be split. Must contain at least one grouping variable.
	Comma-separated list of unquoted variable names that will be used to split the data.
keep_factors	Should the grouping columns be kept?
Х	An object to check for class split_factors.

Details

- split_factors() Split a data frame by factors.
- as.splict_factors() coerce to an object of class split_factors
- is.splict_factors() check if an object is of class split_factors

Value

A list where each element is a named level of the grouping factors. If more than one grouping variable is used, then each element is the combination of the grouping variables.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
g1 <- split_factors(iris, Species)
g2 <- split_factors(data_ge, ENV, keep_factors = TRUE)
spdata <- as.split_factors(iris)
is.split_factors(spdata)</pre>
```

stars_pval

Generate significance stars from p-values

Description

Generate significance stars from p-values using $R\sp{'}s$ standard definitions.

Usage

```
stars_pval(p_value)
```

Arguments

p_value

A numeric vector of p-values

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Details

Mapping from p_value ranges to symbols:

```
0 - 0.0001: '****'
0.0001 - 0.001: '***'
0.001 - 0.05: '*'
0.05 - 1.0: 'ns'
```

Value

A character vector containing the same number of elements as p-value, with an attribute "legend" providing the conversion pattern.

Author(s)

```
Tiago Olivoto <tiagoolivoto@gmail.com>
```

Examples

```
p_{vals} \leftarrow c(0.01, 0.043, 0.1, 0.0023, 0.000012)

stars_{pval}(p_{vals})
```

superiority

Lin e Binns' superiority index

Description

[Stable]

Nonparametric stability analysis using the superiority index proposed by Lin & Binns (1988).

Usage

```
superiority(.data, env, gen, resp, verbose = TRUE)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s)
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
resp	The response variable(s). To analyze multiple variables in a single procedure use, for example, resp = $c(var1, var2, var3)$.
verbose	Logical argument. If verbose = FALSE the code will run silently.

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Value

An object of class superiority where each element is the result of one variable and contains the following items:

- **environments** The mean for each environment, the environment index and classification as favorable and unfavorable environments.
- index The superiority index computed for all (Pi_a), favorable (Pi_f) and unfavorable (Pi_u) environments.

Author(s)

Tiago Olivoto, <tiagoolivoto@gmail.com>

References

Lin, C.S., and M.R. Binns. 1988. A superiority measure of cultivar performance for cultivar x location data. Can. J. Plant Sci. 68:193-198. doi: 10.4141/cjps88018

See Also

```
Annicchiarico(), ecovalence(), ge_stats()
```

Examples

```
library(metan)
out <- superiority(data_ge2, ENV, GEN, PH)
print(out)</pre>
```

themes

Personalized theme for ggplot2-based graphics

Description

- theme_metan(): Theme with a gray background and major grids.
- theme_metan_minimal(): A minimalistic theme with half-open frame, white background, and no grid. For more details see ggplot2::theme().
- transparent_color(): A helper function to return a transparent color with Hex value of "#000000FF"
- ggplot_color(): A helper function to emulate ggplot2 default color palette.
- alpha_color(): Return a semi-transparent color based on a color name and an alpha value. For more details see grDevices::colors().

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Usage

```
theme_metan(grid = "none", col.grid = "white", color.background = "gray95")
theme_metan_minimal()
transparent_color()
ggplot_color(n)
alpha_color(color, alpha = 50)
```

Arguments

grid Control the grid lines in plot. Defaults to "both" (x and y major grids). Allows

also grid = "x" for grids in x axis only, grid = "y" for grid in y axis only, or

grid = "none" for no grids.

col.grid The color for the grid lines

color.background

The color for the panel background.

n The number of colors. This works well for up to about eight colours, but after

that it becomes hard to tell the different colours apart.

color A color name.

alpha An alpha value for transparency (0 < alpha < 1).

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Thennarasu's stability statistics

Description

[Stable]

Performs a stability analysis based on Thennarasu (1995) statistics.

Usage

```
Thennarasu(.data, env, gen, resp, verbose = TRUE)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
resp	The response variable(s). To analyze multiple variables in a single procedure use, for example, $resp = c(var1, var2, var3)$.
verbose	Logical argument. If verbose = FALSE the code will run silently.

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Value

An object of class Thennarasu, which is a list containing the results for each variable used in the argument resp. For each variable, a tibble with the columns GEN, N1, N2, N3 and N4 is returned.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

References

Thennarasu, K. 1995. On certain nonparametric procedures for studying genotype x environment interactions and yield stability. Ph.D. thesis. P.J. School, IARI, New Delhi, India.

Examples

```
library(metan)
out <- Thennarasu(data_ge, ENV, GEN, GY)
print(out)</pre>
```

transpose_df

Transpose a data frame

Description

[Experimental]

Is an alternative to t() to transpose a data frame. The first column of df will become column names in the transposed data.

Usage

```
transpose_df(df)
```

Arguments

df

A data frame to be transposed.

Value

A tibble containing the transposed data.

```
library(metan)
df <-
data.frame(
  GEN = c("G1", "G2", "G3","G4"),
  E1 = rnorm(4, 100, 20),
  E2 = rnorm(4, 10, 2),
  E3 = rnorm(4, 50, 5),
  E4 = rnorm(4, 1000, 150)
)</pre>
```

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```
df
t(df)
transpose_df(df)
```

tukey_hsd

Tukey Honest Significant Differences

Description

[Experimental]

Helper function to perform Tukey post-hoc tests. It is used in gafem.

Usage

```
tukey_hsd(model, ..., out = "long")
```

Arguments

model

an object of class aov or 1m.

. . .

other arguments passed to the function stats::TukeyHSD(). These include:

- which: A character vector listing terms in the fitted model for which the intervals should be calculated. Defaults to all the terms.
- **ordered**: A logical value indicating if the levels of the factor should be ordered according to increasing average in the sample before taking differences. If ordered is true then the calculated differences in the means will all be positive. The significant differences will be those for which the lwr end point is positive.

out

The format of outputs. If out = "long" a 'long' format (tibble) is returned. If out = "wide", a matrix with the adjusted p-values for each term is returned.

Value

A tibble data frame containing the results of the pairwise comparisons (if out = "long") or a "list-columns" with p-values for each term (if out = "wide").

```
library(metan)
mod <- lm(PH ~ GEN + REP, data = data_g)
tukey_hsd(mod)
tukey_hsd(mod, out = "wide")</pre>
```

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utils_as

Encode variables to a specific format

Description

[Stable]

Function to quick encode vector or columns to a specific format.

```
• as_numeric(): Encode columns to numeric using as.numeric().
```

- as_integer(): Encode columns to integer using as.integer().
- as_logical(): Encode columns to logical using as.logical().
- as_character(): Encode columns to character using as.character().
- as_factor(): Encode columns to factor using as.factor().

Usage

```
as_numeric(.data, ..., .keep = "all", .pull = FALSE)
as_integer(.data, ..., .keep = "all", .pull = FALSE)
as_logical(.data, ..., .keep = "all", .pull = FALSE)
as_character(.data, ..., .keep = "all", .pull = FALSE)
as_factor(.data, ..., .keep = "all", .pull = FALSE)
```

Arguments

.data A data frame or a vector.

... <tidy-select>. If .data is a data frame, then ... are the variable(s) to encode to a format.

.keep Allows you to control which columns from .data are retained in the output.

- "all" (default) retains all variables.
- "used" keeps any variables used to make new variables.

.pull Allows you to pull out the last column of the output. It is useful in combination with .keep = "used". In this case, a vector will be created with the used column.

Value

An object of the same class of .data with the variables in ... encoded to the specified format.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

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Examples

```
library(metan)
library(tibble)
df <-
  tibble(y = rnorm(5),
         x1 = c(1:5),
         x2 = c(TRUE, TRUE, FALSE, FALSE, FALSE),
         x3 = letters[1:5],
         x4 = as.factor(x3))
df
# Convert y to integer
as_integer(df, y)
as_integer(df$y)
# convert x3 to factor
as_factor(df, x3)
# Convert all columns to character
as_character(df, everything())
# Convert x2 to numeric and coerce to a vector
as_numeric(df, x2, .keep = "used", .pull = TRUE)
```

utils_bind

Helper function for binding data

Description

- rbind_fill() [Deprecated] use rbind_fill_id() instead.
- rbind_fill_id() [Experimental] Implements the common pattern of do.call(rbind, dfs) with data frame identifier and filling of missing values.

Usage

```
rbind_fill_id(..., .id = NULL, .fill = NA)
rbind_fill(..., .fill = NA)
```

Arguments

.id

... The dataframes. Either a list of data frames, or a comma-separated list of dataframes.

Data frame identifier. If a comma-separated list of data frames is supplied, the labels are taken from the names of the objects. When a list of data frames is supplied, the labels are taken from the names of the list. If no names are found, a numeric sequence is used instead.

.fill When row-binding, columns are matched by name, and any missing columns will be filled with NA Defaults to NA.

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Value

A data frame.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

utils_class

Utilities for handling with classes

Description

Utilities for handling with classes

Usage

```
add_class(x, class)
has_class(x, class)
remove_class(x, class)
set_class(x, class)
```

Arguments

x An object

class The class to add or remove

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Details

- add_class(): add a class to the object x keeping all the other class(es).
- has_class(): Check if a class exists in object x and returns a logical value.
- set_class(): set a class to the object x.
- remove_class(): remove a class from the object x.

Value

The object x with the class added or removed.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

Examples

```
library(metan)
df <-
data_ge2 %>%
add_class("my_class")
class(df)
has_class(df, "my_class")
remove_class(df, "my_class") %>% class()
set_class(df, "data_frame") %>% class()
```

utils_data

Utilities for data Copy-Pasta

Description

[Stable]

These functions allows interacting with the system clipboard. It is possible read from the clipboard or write a data frame or matrix to the clipboard.

- clip_read() read data from the clipboard.
- clip_write() write data to the clipboard.

Usage

```
clip_read(header = TRUE, sep = "\t", ...)
clip_write(.data, sep = "\t", row_names = FALSE, col_names = TRUE, ...)
```

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Arguments

header	If the copied data has a header row for dataFrame, defaults to TRUE.
sep	The separator which should be used in the copied output.
	Further arguments to be passed to utils::read.table().
.data	The data that should be copied to the clipboard. Only data frames and matrices are allowed
row_names	Decides if the output should keep row names or not, defaults to FALSE.
col_names	Decides if the output should keep column names or not, defaults to TRUE.

Value

Nothing

Author(s)

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utils_data_org Utilities for data organization
--

Description

[Experimental]

Useful function for data organization before statistical analysis

- add_seq_block(): Add a column with sequential block numeration in multi-environment data sets.
- recode_factor(): Recode a factor column. A sequential numbering (with possible prefix) is used to identify each level.
- df_to_selegen_54(): Given a multi-environment data with environment, genotype, and replication, format the data to be used in the Selegen software (model 54).

Usage

```
add_seq_block(data, env, rep, new_factor = BLOCK, prefix = "", verbose = TRUE)
recode_factor(data, factor, new_factor = CODE, prefix = "", verbose = TRUE)
df_to_selegen_54(data, env, gen, rep, verbose = TRUE)
```

Arguments

data	A data frame.
env	The name of the column that contains the levels of the environments.
rep	The name of the column that contains the levels of the replications/blocks.
new_factor	The name of the new column created.
prefix	An optional prefix to bind with the new factor.

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verbose Logical argument. If verbose = FALSE the code will run silently.

factor A column to recode.

gen The name of the column that contains the levels of the genotypes, that will be

treated as random effect.

Author(s)

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References

Resende, M.D. V. 2016. Software Selegen-REML/BLUP: a useful tool for plant breeding. Crop Breed. Appl. Biotechnol. 16(4): 330–339. doi: 10.1590/198470332016v16n4a49.

Examples

```
library(metan)
df_ge <- ge_simula(ngen = 2,</pre>
                   nenv = 3,
                   nrep = 2) %>%
         add_cols(ENV = c(rep("CACIQUE", 4),
                          rep("FREDERICO", 4),
                          rep("SANTA_MARIA", 4)))
df_ge
# Add sequential block numbering over environments
add_seq_block(df_ge, ENV, REP, prefix = "B")
# Recode the 'ENV' column to "ENV1", "ENV2", and so on.
recode_factor(df_ge,
              factor = ENV,
              prefix = "ENV",
              new_factor = ENV_CODE)
# Format the data to be used in the Selegen software (model 54)
df <- df_to_selegen_54(df_ge, ENV, GEN, REP) %>%
recode_factor(ENV, prefix = "E", new_factor = ENV)
```

utils_mat

Utilities for handling with matrices

Description

[Stable]

These functions help users to make upper, lower, or symmetric matrices easily.

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Usage

```
make_upper_tri(x, diag = NA)
make_lower_tri(x, diag = NA)
make_lower_upper(lower, upper, diag = NA)
make_sym(x, make = "upper", diag = NA)
tidy_sym(x, keep_diag = TRUE)
```

Arguments

X	A matrix to apply the function. It must be a symmetric (square) matrix in make_upper_tri() and make_lower_tri() or a triangular matrix in make_sym(). tidy_sym() accepts both symmetrical or triangular matrices.
diag	What show in the diagonal of the matrix. Default to NA.
lower	A square matrix to fill the lower diagonal of the new matrix.
upper	A square matrix to fill the upper diagonal of the new matrix.
make	The triangular to built. Default is "upper". In this case, a symmetric matrix will be built based on the values of a lower triangular matrix.
keep_diag	Keep diagonal values in the tidy data frame? Defaults to TRUE.

Details

- make_upper_tri() makes an upper triangular matrix using a symmetric matrix.
- make_lower_tri() makes a lower triangular matrix using a symmetric matrix.
- make_sym() makes a lower triangular matrix using a symmetric matrix.
- tidy_sym() transform a symmetric matrix into tidy data frame.

Value

An upper, lower, or symmetric matrix, or a tidy data frame.

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```
library(metan)
m <- cor(select_cols(data_ge2, 5:10))
make_upper_tri(m)
make_lower_tri(m)
make_lower_tri(m) %>%
make_sym(diag = 0)
tidy_sym(m)
tidy_sym(make_lower_tri(m))
```

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utils_na_zero

Utilities for handling with NA and zero values

Description

[Stable]

NAs and zeros can increase the noise in multi-environment trial analysis. This collection of functions will make it easier to deal with them.

- fill_na(): Fills NA in selected columns using the next or previous entry.
- has_na(), has_zero(): Check for NAs and 0s in the data and return a logical value.
- random_na(): Generate random NA values in a two-way table based on a desired proportion.
- remove_cols_na(), remove_cols_zero(): Remove columns with NAs and 0s, respectively.
- remove_rows_na(), remove_rows_zero(): Remove rows with NAs and 0s, respectively.
- select_cols_na(), select_cols_zero(): Select columns with NAs and 0s, respectively.
- select_rows_na(), select_rows_zero(): Select rows with NAs and Os, respectively.
- replace_na(), replace_zero(): Replace NAs and Os, respectively, with a replacement value.

Usage

```
fill_na(.data, ..., direction = "down")
has_na(.data)

remove_rows_na(.data, verbose = TRUE)

remove_cols_na(.data, verbose = TRUE)

select_cols_na(.data, verbose = TRUE)

select_rows_na(.data, verbose = TRUE)

replace_na(.data, ..., replacement = 0)

random_na(.data, prop)
has_zero(.data)

remove_rows_zero(.data, verbose = TRUE)

remove_cols_zero(.data, verbose = TRUE)

select_cols_zero(.data, verbose = TRUE)

select_rows_zero(.data, verbose = TRUE)

replace_zero(.data, ..., replacement = NA)
```

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Arguments

.data	A data frame.
	Variables to fill NAs in fill_na(), replace NAs in replace_na() or zeros in replace_zero(). If is null then all variables in .data will be evaluated. It must be a single variable name or a comma-separated list of unquoted variables names. Select helpers are also allowed.
direction	Direction in which to fill missing values. Currently either "down" (the default), "up", "downup" (i.e. first down and then up) or "updown" (first up and then down).
verbose	Logical argument. If TRUE (default) shows in console the rows or columns deleted.
replacement	The value used for replacement. Defaults to 0. Use replacement. = "colmean" to replace missing values with column mean.
prop	The proportion (percentage) of NA values to generate in .data.

Value

A data frame with rows or columns with NA values deleted.

Author(s)

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```
library(metan)
data_naz <- iris %>%
              group_by(Species) %>%
              doo(\sim head(., n = 3)) \%>\%
              as_character(Species)
data_naz
data_naz[c(2:3, 6, 8), c(1:2, 4, 5)] \leftarrow NA
data_naz[c(2, 7, 9), c(2, 3, 4)] <- 0
has_na(data_naz)
has_zero(data_naz)
\mbox{\# Fill NA} values of column GEN
fill_na(data_naz, Species)
# Remove columns
remove_cols_na(data_naz)
remove_cols_zero(data_naz)
remove_rows_na(data_naz)
remove_rows_zero(data_naz)
# Select columns
select_cols_na(data_naz)
select_cols_zero(data_naz)
select_rows_na(data_naz)
select_rows_zero(data_naz)
# Replace values
replace_na(data_naz)
```

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```
replace_zero(data_naz)
```

utils_num_str

Utilities for handling with numbers and strings

Description

[Stable]

- all_lower_case(): Translate all non-numeric strings of a data frame to lower case.
- all_upper_case(): Translate all non-numeric strings of a data frame to upper case.
- all_title_case(): Translate all non-numeric strings of a data frame to title case.
- first_upper_case: Translate the first word of a string to upper case.
- extract_number(): Extract the number(s) of a string.
- extract_string(): Extract all strings, ignoring case.
- find_text_in_num(): Find text characters in a numeric sequence and return the row index.
- has_text_in_num(): Inspect columns looking for text in numeric sequence and return a warning if text is found.
- remove_space(): Remove all blank spaces of a string.
- remove_strings(): Remove all strings of a variable.
- replace_number(): Replace numbers with a replacement.
- replace_string(): Replace all strings with a replacement, ignoring case.
- round_cols(): Round a selected column or a whole data frame to significant figures.
- tidy_strings(): Tidy up characters strings, non-numeric columns, or any selected columns in a data frame by putting all word in upper case, replacing any space, tabulation, punctuation characters by '_', and putting '_' between lower and upper case. Suppose that str = c("Env1", "env 1", "env.1") (which by definition should represent a unique level in plant breeding trials, e.g., environment 1) is subjected to tidy_strings(str): the result will be then c("ENV_1", "ENV_1", "ENV_1"). See Examples section for more examples.

Usage

```
all_upper_case(.data, ...)
all_lower_case(.data, ...)
all_title_case(.data, ...)
first_upper_case(.data, ...)
extract_number(.data, ..., pattern = NULL)
extract_string(.data, ..., pattern = NULL)
find_text_in_num(.data, ...)
```

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```
has_text_in_num(.data)
remove_space(.data, ...)
remove_strings(.data, ...)
replace_number(
  .data,
  . . . ,
  pattern = NULL,
  replacement = ""
  ignore_case = FALSE
replace_string(
  .data,
  pattern = NULL,
  replacement = "",
  ignore_case = FALSE
round_cols(.data, ..., digits = 2)
tidy_strings(.data, ..., sep = "_")
```

Arguments

.data A data frame

... The argument depends on the function used.

- For round_cols() ... are the variables to round. If no variable is informed, all the numeric variables from data are used.
- For all_lower_case(), all_upper_case(), all_title_case(), stract_number(), stract_string(), remove_strings(), and tidy_strings() ... are the variables to apply the function. If no variable is informed, the function will be applied to all non-numeric variables in .data.

pattern A string to be matched. Regular Expression Syntax is also allowed.

replacement A string for replacement.

ignore_case If FALSE (default), the pattern matching is case sensitive and if TRUE, case is

ignored during matching.

digits The number of significant figures.

sep A character string to separate the terms. Defaults to "_".

Author(s)

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```
library(metan)
```

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```
# All numeric columns
round_cols(data_ge2, digits = 1)
# Round specific columns
round_cols(data_ge2, EP, digits = 1)
######## Extract or replace numbers ########
# Extract numbers
extract_number(data_ge, GEN)
# Replace numbers
replace_number(data_ge, GEN)
replace_number(data_ge,
             GEN,
             pattern = 1,
             replacement = "_one")
####### Extract, replace or remove strings ########
# Extract strings
extract_string(data_ge, GEN)
# Replace strings
replace_string(data_ge, GEN)
replace_string(data_ge,
             GEN,
             pattern = "G",
             replacement = "GENOTYPE_")
# Remove strings
remove_strings(data_ge)
remove_strings(data_ge, ENV)
######### Find text in numeric sequences #########
mixed_text <- data.frame(data_ge)</pre>
mixed_text[2, 4] <- "2..503"
mixed_text[3, 4] <- "3.2o75"
find_text_in_num(mixed_text, GY)
######### upper, lower and title cases #########
gen_text <- c("This is the first string.", "this is the second one")</pre>
all_lower_case(gen_text)
all_upper_case(gen_text)
all_title_case(gen_text)
first_upper_case(gen_text)
# A whole data frame
all_lower_case(data_ge)
messy_env <- c("ENV 1", "Env 1", "Env1", "env1", "Env.1", "Env_1")
tidy_strings(messy_env)
messy_gen <- c("GEN1", "gen 2", "Gen.3", "gen-4", "Gen_5", "GEN_6")
tidy_strings(messy_gen)
```

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utils_progress

Utilities for text progress bar in the terminal

Description

[Experimental]

Progress bars are configurable, may include percentage, elapsed time, and custom text.

- progress(): Initiate a custom progress bar of class pb_metan.
- run_progress(): Run the progress bar and should be called within a 'for loop' statement, a lapply() family or purrr::map() family of functional programming tools.

Usage

```
progress(
    min = 0,
    max = 100,
    leftd = "|",
    rightd = "|",
    char = "=",
    style = 2,
    width = getOption("width"),
    time = Sys.time()
)
run_progress(pb, actual, text = "", digits = 0, sleep = 0)
```

Arguments

min, max

Numeric values for the extremes of the progress bar. Must have min < max.

leftd, rightd

The left and right delimiters for the progress bar. Defaults to "|".

that

The character (or character string) to form the progress bar.

style

The 'style' of the progress bar. Elapsed time is counted from calling progress() up to each call of run_progress().

- type = 1: Shows a progress bar without percentage or elapsed time.
- type = 2: The default, shows the progress bar and its percentage.

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•	type =	3:	Shows	the	progress	bar	and	elapsed	l time.
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• type = 4: Shows the progress bar, percentage, and 6	elapsed time.
---	---------------

width The the width of the progress bar. Defaults to the number of characters is that which fits into getOption("width"). time The system time used to compute the elapsed time from calling progress() to each call of run_progress(). Defaults to Sys.time(). An object created with progress() pb actual The actual value, for example, a loop variable that define the loop index value. An optional character string to be shown at the begining of the progress bar. text digits The number of significant figures in percentage value. Defaults to 0. Suspend execution for a time interval with Sys.sleep() within run_progress(). sleep

Defaults to 0.

Value

progress() returns a list of class pb_metan that contains the set parameters that will called by run_progress().

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```
pb <- progress()</pre>
for (i in 1:100) {
run_progress(pb, actual = i, sleep = 0.01)
############## An apply family approach ############
pb <- progress(max = 10)</pre>
foo <- function(...){</pre>
   run_progress(pb, ...)
   rnorm(100) %>% mean()
}
(a <- sapply(1:10, FUN = foo, sleep = 0.05))
####### A purrr functional programming approach ########
foo2 <- function(...){</pre>
    run_progress(pb2, ...)
    rnorm(100) %>% mean()
pb2 \leftarrow progress(max = 10000,
               style = 4,
               leftd = "",
               char = "."
               rightd = "!")
b <- purrr::map_dbl(1:10000, foo2, text = "Progress bar for sampling")</pre>
hist(b)
```

utils_rows_cols

Utilities for handling with rows and columns

Description

[Stable]

- add_cols(): Add one or more columns to an existing data frame. If specified .before or .after columns does not exist, columns are appended at the end of the data. Return a data frame with all the original columns in .data plus the columns declared in In add_cols() columns in .data are available for the expressions. So, it is possible to add a column based on existing data.
- add_rows(): Add one or more rows to an existing data frame. If specified .before or .after rows does not exist, rows are appended at the end of the data. Return a data frame with all the original rows in .data plus the rows declared in . . . argument.
- add_row_id(): Add a column with the row id as the first column in .data.
- add_prefix() and add_suffix() add prefixes and suffixes, respectively, in variable names selected in . . . argument.
- all_pairs(): Get all the possible pairs between the levels of a factor.
- colnames_to_lower(): Translate all column names to lower case.
- colnames_to_upper(): Translate all column names to upper case.
- colnames_to_title(): Translate all column names to title case.
- column_exists(): Checks if a column exists in a data frame. Return a logical value.
- columns_to_first(): Move columns to first positions in .data.
- columns_to_last(): Move columns to last positions in .data.
- columns_to_rownames(): Move a column of .data to its row names.
- rownames_to_column(): Move the row names of .data to a new column.
- remove_rownames(): Remove the row names of .data.
- concatenate(): Concatenate columns of a data frame. If drop = TRUE then the existing variables are dropped. If pull = TRUE then the concatenated variable is pull out to a vector. This is specially useful when using concatenate to add columns to a data frame with add_cols().
- get_levels(): Get the levels of a factor variable.
- get_levels_comb(): Get the combination of the levels of a factor.
- get_level_size(): Get the size of each level of a factor variable.
- remove_cols(): Remove one or more columns from a data frame.
- remove_rows(): Remove one or more rows from a data frame.
- reorder_cols(): Reorder columns in a data frame.
- select_cols(): Select one or more columns from a data frame.
- select_first_col(): Select first variable, possibly with an offset.
- select_last_col(): Select last variable, possibly with an offset.
- select_numeric_cols(): Select all the numeric columns of a data frame.
- select_non_numeric_cols(): Select all the non-numeric columns of a data frame.
- select_rows(): Select one or more rows from a data frame.
- tidy_colnames(): Tidy up column names with tidy_strings().

Usage

```
add_cols(.data, ..., .before = NULL, .after = NULL)
add_rows(.data, ..., .before = NULL, .after = NULL)
add_row_id(.data, var = "row_id")
all_pairs(.data, levels)
add_prefix(.data, ..., prefix, sep = "_")
add_suffix(.data, ..., suffix, sep = "_")
colnames_to_lower(.data)
colnames_to_upper(.data)
colnames_to_title(.data)
column_to_first(.data, ...)
column_to_last(.data, ...)
column_to_rownames(.data, var = "rowname")
rownames_to_column(.data, var = "rowname")
remove_rownames(.data, ...)
column_exists(.data, cols)
concatenate(
  .data,
  ...,
 prefix = NULL,
  suffix = NULL,
 new_var = new_var,
 sep = "_",
 drop = FALSE,
 pull = FALSE,
  .before = NULL,
  .after = NULL
get_levels(.data, ...)
get_levels_comb(.data, ...)
get_level_size(.data, ...)
reorder_cols(.data, ..., .before = NULL, .after = NULL)
```

```
remove_cols(.data, ...)
remove_rows(.data, ...)
select_first_col(.data, offset = NULL)
select_last_col(.data, offset = NULL)
select_numeric_cols(.data)
select_non_numeric_cols(.data)
select_cols(.data, ...)
select_rows(.data, ...)
tidy_colnames(.data, sep = "_")
```

Arguments

.data A data frame

... The argument depends on the function used.

• For add_cols() and add_rows() is name-value pairs. All values must have one element for each row in .data when using add_cols() or one element for each column in .data when using add_rows(). Values of length 1 will be recycled when using add_cols().

- For remove_cols() and select_cols(), ... is the column name or column index of the variable(s) to be dropped.
- For add_prefix() and add_suffix(), ... is the column name to add the prefix or suffix, respectively. Select helpers are allowed.
- For columns_to_first() and columns_to_last(), ... is the column name or column index of the variable(s) to be moved to first or last in .data.
- For remove_rows() and select_rows(), ... is an integer row value.
- For concatenate(), ... is the unquoted variable names to be concatenated
- For get_levels(), get_level_comb(), and get_level_size() ... is the unquoted variable names to get the levels, levels combinations and levels size, respectively.

.before, .after

For add_cols(), concatenate(), and reorder_cols(), one-based column index or column name where to add the new columns, default: .after last column. For add_rows(), one-based row index where to add the new rows, default: .after last row

var Name of column to use for rownames.

levels The levels of a factor or a numeric vector.

prefix, suffix The prefix and suffix used in add_prefix() and add_suffix(), respectively.

sep The separator to appear when using concatenate(), add_prefix(), or add_suffix().

Defaults to to "_".

cols A quoted variable name to check if it exists in .data.

new_var	The name of the new variable containing the concatenated values. Defaults to new_var.
drop	Logical argument. If TRUE keeps the new variable new_var and drops the existing ones. Defaults to FALSE.
pull	Logical argument. If TRUE, returns the last column (on the assumption that's the column you've created most recently), as a vector.
offset	Set it to <i>n</i> to select the <i>n</i> th variable from the end (for select_last_col()) of from the begin (for select_first_col())

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```
library(metan)
\mbox{\tt\#} Variables x and y .after last column
data_ge %>%
 add_cols(x = 10,
         y = 30)
\# Variables x and y .before the variable GEN
data_ge %>%
  add_cols(x = 10,
         y = 30,
          .before = GEN)
# Creating a new variable based on the existing ones.
data_ge %>%
  add_cols(GY2 = GY^2,
          GY2_HM = GY2 + HM,
          .after = GY)
reorder_cols(data_ge2, NKR, .before = ENV)
reorder_cols(data_ge2, where(is.factor), .after = last_col())
####### Selecting and removing columns ########
select_cols(data_ge2, GEN, REP)
remove_cols(data_ge2, GEN, REP)
####### Selecting and removing rows #########
select_rows(data_ge2, 2:3)
remove_rows(data_ge2, 2:3)
######## Concatenating columns ###############
{\tt concatenate(data\_ge,\ ENV,\ GEN,\ REP)}
concatenate(data_ge, ENV, GEN, REP, drop = TRUE)
# Combine with add_cols() and replace_string()
data_ge2 %>%
add_cols(ENV_GEN = concatenate(., ENV, GEN, pull = TRUE),
         .after = GEN) %>%
 replace_string(ENV_GEN,
```

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```
pattern = "H",
               replacement = "HYB_")
# Use prefixes and suffixes
concatenate(data_ge2, REP, prefix = "REP", new_var = REP)
# Use prefixes and suffixes (the ear traits EH, EP, EL, and ED)
add_prefix(data_ge2, PH, EH, EP, EL, prefix = "EAR")
add_suffix(data_ge2, PH, EH, EP, EL, suffix = "EAR", sep = ".")
# Use prefixes and suffixes (colnames)
concatenate(data_ge2, REP, prefix = "REP", new_var = REP)
######## formating column names #############
# Creating data with messy column names
df <- head(data_ge, 3)</pre>
colnames(df) <- c("Env", "gen", "Rep", "GY", "hm")</pre>
colnames_to_lower(df)
colnames_to_upper(df)
colnames_to_title(df)
data_ge %>%
  add_rows(GY = 10.3,
          HM = 100.11,
          .after = 1)
######## checking if a column exists #########
column_exists(data_g, "GEN")
###### get the levels, level combinations and size of levels #######
get_levels(data_g, GEN)
get_levels_comb(data_ge, ENV, GEN)
get_level_size(data_g, GEN)
########### all possible pairs ###############
all_pairs(data_g, GEN)
######## select numeric variables only ########
select_numeric_cols(data_g)
select_non_numeric_cols(data_g)
```

utils_sets

Utilities for set operations for many sets

Description

[Stable]

Provides alternative function to base::union(), base::intersect(), and base::setdiff().

• set_union(): Returns the union of the sets in

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- set_intersect(): Returns the intersect of the sets in
- set_difference(): Returns the difference of the sets in

Usage

```
set_intersect(..., pairs = FALSE)
set_union(..., pairs = FALSE)
set_difference(..., pairs = FALSE)
```

Arguments

. . .

A list or a comma-separated list of vectors in the same class. If vector contains duplicates they will be discarded. If the list doesn't have names the sets will be named as "set_1", "Set_2", "Set_3" and so on. If vectors are given in . . . , the set names will be named with the names of the objects provided. Data frames are also allowed, provided that common column names exist.

pairs

Returns the pairwise unions of the sets? Defaults to FALSE.

Value

A vector showing the desired operation of the sets. If pairs = TRUE, returns a list showing the pairwise operation of the sets.

Author(s)

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```
library(metan)
(A <- letters[1:4])
(B <- letters[2:5])
(C <- letters[3:7])
set_union(A, B)
set_intersect(A, B, C)
set_difference(B, C)
# Operations with data frames
# Add a row id for better understanding
sets <- data_ge %>% add_row_id()
set_1 <- sets[1:5,]
set_2 <- sets[2:6,]
set_3 <- sets[3:7,]
set_intersect(set_1, set_2, set_3)
set_difference(set_2, set_3)
set_union(set_1, set_2, set_3)
```

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utils_stats

Useful functions for computing descriptive statistics

Description

[Stable]

- The following functions compute descriptive statistics by levels of a factor or combination of factors quickly.
 - cv_by() For computing coefficient of variation.
 - max_by() For computing maximum values.
 - means_by() For computing arithmetic means.
 - min_by() For compuing minimum values.
 - n_by() For getting the length.
 - sd_by() For computing sample standard deviation.
 - sem_by() For computing standard error of the mean.
- Useful functions for descriptive statistics. All of them work naturally with \%>\%, handle grouped data and multiple variables (all numeric variables from .data by default).
 - av_dev() computes the average absolute deviation.
 - ci_mean() computes the confidence interval for the mean.
 - cv() computes the coefficient of variation.
 - freq_table() Computes frequency fable. Handles grouped data.
 - hmean(), gmean() computes the harmonic and geometric means, respectively. The harmonic mean is the reciprocal of the arithmetic mean of the reciprocals. The geometric mean is the *n*th root of *n* products.
 - kurt() computes the kurtosis like used in SAS and SPSS.
 - range_data() Computes the range of the values.
 - n_valid() The valid (not NA) length of a data.
 - n_unique() Number of unique values.
 - n_missing() Number of missing values.
 - row_col_mean(), row_col_sum() Adds a row with the mean/sum of each variable and a column with the mean/sum for each row of the data.
 - sd_amo(), sd_pop() Computes sample and populational standard deviation, respectively.
 - sem() computes the standard error of the mean.
 - skew() computes the skewness like used in SAS and SPSS.
 - sum_dev() computes the sum of the absolute deviations.
 - sum_sq() computes the sum of the squared values.
 - sum_sq_dev() computes the sum of the squared deviations.
 - var_amo(), var_pop() computes sample and populational variance.

desc_stat() is wrapper function around the above ones and can be used to compute quickly all
these statistics at once.

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Usage

```
av_dev(.data, ..., na.rm = FALSE)
ci_mean(.data, ..., na.rm = FALSE, level = 0.95)
cv(.data, ..., na.rm = FALSE)
freq_table(.data, ...)
hmean(.data, ..., na.rm = FALSE)
gmean(.data, ..., na.rm = FALSE)
kurt(.data, ..., na.rm = FALSE)
n_missing(.data, ..., na.rm = FALSE)
n_unique(.data, ..., na.rm = FALSE)
n_valid(.data, ..., na.rm = FALSE)
pseudo_sigma(.data, ..., na.rm = FALSE)
range_data(.data, ..., na.rm = FALSE)
row_col_mean(.data, na.rm = FALSE)
row_col_sum(.data, na.rm = FALSE)
sd_amo(.data, ..., na.rm = FALSE)
sd_pop(.data, ..., na.rm = FALSE)
sem(.data, ..., na.rm = FALSE)
skew(.data, ..., na.rm = FALSE)
sum_dev(.data, ..., na.rm = FALSE)
sum_sq_dev(.data, ..., na.rm = FALSE)
sum_sq(.data, ..., na.rm = FALSE)
var_pop(.data, ..., na.rm = FALSE)
var_amo(.data, ..., na.rm = FALSE)
cv_by(.data, ..., na.rm = FALSE)
max_by(.data, ..., na.rm = FALSE)
means_by(.data, ..., na.rm = FALSE)
```

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```
min_by(.data, ..., na.rm = FALSE)
n_by(.data, ..., na.rm = FALSE)
sd_by(.data, ..., na.rm = FALSE)
sem_by(.data, ..., na.rm = FALSE)
sum_by(.data, ..., na.rm = FALSE)
```

Arguments

.data A data frame or a numeric vector.

... The argument depends on the function used.

- For *_by functions, ... is one or more categorical variables for grouping the data. Then the statistic required will be computed for all numeric variables in the data. If no variables are informed in ..., the statistic will be computed ignoring all non-numeric variables in .data.
- For the other statistics, ... is a comma-separated of unquoted variable names to compute the statistics. If no variables are informed in n ..., the statistic will be computed for all numeric variables in .data.

na.rm If FALSE, the default, missing values are removed with a warning. If TRUE, missing values are silently removed.

level The confidence level for the confidence interval of the mean. Defaults to 0.95.

Value

- Functions *_by() returns a tbl_df with the computed statistics by each level of the factor(s) declared in
- All other functions return a nammed integer if the input is a data frame or a numeric value if the input is a numeric vector.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

```
library(metan)
# means of all numeric variables by ENV
means_by(data_ge2, GEN, ENV)

# Coefficient of variation for all numeric variables
# by GEN and ENV
cv_by(data_ge2, GEN, ENV)

# Skewness of a numeric vector
set.seed(1)
nvec <- rnorm(200, 10, 1)
skew(nvec)

# Confidence interval 0.95 for the mean</pre>
```

venn_plot

```
# All numeric variables
# Grouped by levels of ENV
data_ge2 %>%
   group_by(ENV) %>%
   ci_mean()

# standard error of the mean
# Variable PH and EH
sem(data_ge2, PH, EH)

# Frequency table for variable NR
data_ge2 %>%
   freq_table(NR)
```

venn_plot

Draw Venn diagrams

Description

[Stable]

Produces ggplot2-based Venn plots for 2, 3 or 4 sets. A Venn diagram shows all possible logical relationships between several sets of data.

Usage

```
venn_plot(
  ...,
  names = NULL,
  show_elements = FALSE,
  show_sets = FALSE,
  fill = ggplot_color(4),
  alpha = 0.5,
  stroke_color = "white",
  stroke_alpha = 1,
  stroke_size = 1,
  stroke_linetype = "solid",
  name_color = "black",
  name\_size = 6,
  text_color = "black",
  text_size = 4,
  label_sep = ","
)
```

Arguments

. . .

A list or a comma-separated list of vectors in the same class. If vector contains duplicates they will be discarded. If the list doesn't have names the sets will be named as "set_1", "Set_2", "Set_3" and so on. If vectors are given in . . . , the set names will be named with the names of the objects provided.

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names By default, the names of the sets are set as the names of the objects in ... (names = NULL). Use names to override this default. show_elements Show set elements instead of count. Defaults to FALSE. show_sets Show set names instead of count. Defaults to FALSE. Filling colors in circles. Defaults to the default ggplot2 color palette. A vector fill of length 1 will be recycled. alpha Transparency for filling circles. Defaults to 0.5. Stroke color for drawing circles. stroke_color stroke_alpha Transparency for drawing circles. stroke_size Stroke size for drawing circles. stroke_linetype Line type for drawing circles. Defaults to "solid". Text color for set names. Defaults to "black". name_color Text size for set names. name_size text_color Text color for intersect contents. text_size Text size for intersect contents. label_sep The separator for labs when show_elements = TRUE. Defaults to ", ".

Value

A ggplot object.

Author(s)

Tiago Olivoto <tiagoolivoto@gmail.com>

```
library(metan)
(A <- letters[1:4])
(B <- letters[2:5])
(C <- letters[3:7])
(D <- letters[4:12])
# create a Venn plot
venn_plot(A, B)
# Three sets
venn_plot(A, B, C)
# Four sets
venn_plot(A, B, C, D)
# Use a list
dfs \leftarrow list(A = A, B = B, C = C, D = D)
venn_plot(dfs,
          show_elements = TRUE,
          fill = c("red", "blue", "green", "gray"),
          stroke_color = "black",
          alpha = 0.8,
```

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```
text_size = 8,
label_sep = ".")
```

waas

Weighted Average of Absolute Scores

Description

[Stable]

Compute the Weighted Average of Absolute Scores for AMMI analysis (Olivoto et al., 2019).

This function compute the weighted average of absolute scores, estimated as follows:

$$WAAS_i = \sum_{k=1}^{p} |IPCA_{ik} \times EP_k| / \sum_{k=1}^{p} EP_k$$

where $WAAS_i$ is the weighted average of absolute scores of the ith genotype; $IPCA_{ik}$ is the score of the ith genotype in the kth IPCA; and EP_k is the explained variance of the kth IPCA for k=1,2,...p, considering p the number of significant PCAs, or a declared number of PCAs. For example if prob = 0.05, all axis that are significant considering this probability level are used. The number of axis can be also informed by declaring naxis = x. This will override the number of significant axes according to the argument codeprob.

Usage

```
waas(
   .data,
   env,
   gen,
   rep,
   resp,
   block = NULL,
   mresp = NULL,
   wresp = NULL,
   prob = 0.05,
   naxis = NULL,
   ind_anova = FALSE,
   verbose = TRUE
)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
rep	The name of the column that contains the levels of the replications/blocks.
resp	The response variable(s). To analyze multiple variables in a single procedure a vector of variables may be used. For example $resp = c(var1, var2, var3)$.

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block Defaults to NULL. In this case, a randomized complete block design is considered. If block is informed, then a resolvable alpha-lattice design (Patterson and

Williams, 1976) is employed. **All effects, except the error, are assumed to be**

fixed.

mresp The new maximum value after rescaling the response variable. By default, all

variables in resp are rescaled so that de maximum value is 100 and the minimum value is 0 (i.e., mresp = NULL). It must be a character vector of the same length of resp if rescaling is assumed to be different across variables, e.g., if for the first variable smaller values are better and for the second one, higher values are better, then mresp = c("1,h") must be used. Character value of length 1 will

be recycled with a warning message.

wresp The weight for the response variable(s) for computing the WAASBY index. By

default, all variables in resp have equal weights for mean performance and stability (i.e., wresp = 50). It must be a numeric vector of the same length of resp to assign different weights across variables, e.g., if for the first variable equal weights for mean performance and stability are assumed and for the second one, a higher weight for mean performance (e.g. 65) is assumed, then wresp = c(50,65) must be used. Numeric value of length 1 will be recycled with a

warning message.

prob The p-value for considering an interaction principal component axis significant.

naxis The number of IPCAs to be used for computing the WAAS index. Default is

NULL (Significant IPCAs are used). If values are informed, the number of IPCAS will be used independently on its significance. Note that if two or more variables

are included in resp, then naxis must be a vector.

ind_anova Logical argument set to FALSE. If TRUE an within-environment ANOVA is per-

formed.

verbose Logical argument. If verbose = FALSE the code is run silently.

Value

An object of class waas with the following items for each variable:

- individual A within-environments ANOVA considering a fixed-effect model.
- model A data frame with the response variable, the scores of all Principal Components, the estimates of Weighted Average of Absolute Scores, and WAASY (the index that consider the weights for stability and productivity in the genotype ranking.
- MeansGxE The means of genotypes in the environments
- PCA Principal Component Analysis.
- anova Joint analysis of variance for the main effects and Principal Component analysis of the interaction effect.
- **Details** A list summarizing the results. The following information are showed. WgtResponse, the weight for the response variable in estimating WAASB, WgtWAAS the weight for stability, Ngen the number of genotypes, Nenv the number of environments, OVmean the overall mean, Min the minimum observed (returning the genotype and environment), Max the maximum observed, Max the maximum observed, MinENV the environment with the lower mean, MaxENV the environment with the larger mean observed, MinGEN the genotype with the lower mean, MaxGEN the genotype with the larger.
- augment: Information about each observation in the dataset. This includes predicted values in the fitted column, residuals in the resid column, standardized residuals in the stdres column, the diagonal of the 'hat' matrix in the hat, and standard errors for the fitted values in the se.fit column.

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• **probint** The p-value for the genotype-vs-environment interaction.

Author(s)

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References

Olivoto, T., A.D.C. L\'ucio, J.A.G. da silva, V.S. Marchioro, V.Q. de Souza, and E. Jost. 2019a. Mean performance and stability in multi-environment trials I: Combining features of AMMI and BLUP techniques. Agron. J. 111:2949-2960. doi: 10.2134/agronj2019.03.0220

See Also

```
waas_means() waasb() get_model_data()
```

```
library(metan)
# Example 1: Analyzing all numeric variables considering p-value#
# <= 0.05 to compute the WAAS.
model <- waas(data_ge,</pre>
         env = ENV,
         gen = GEN,
         rep = REP,
         resp = everything())
# Residual plot (first variable)
plot(model)
# Get the WAAS index
get_model_data(model, "WAAS")
# Plot WAAS and response variable
plot_scores(model, type = 3)
# Example 2: Declaring the number of axis to be used for
\# computing WAAS and assigning a larger weight for the response \#
# variable when computing the WAASBY index.
model2 <- waas(data_ge,</pre>
           env = ENV,
           gen = GEN,
           rep = REP,
           resp = everything(),
           naxis = 1, # Only to compare with PC1
           wresp = 60)
# Get the WAAS index (it will be |PC1|)
get_model_data(model2)
# Get values for IPCA1
get_model_data(model2, "PC1")
```

waasb

Weighted Average of Absolute Scores

Description

[Stable]

Compute the Weighted Average of Absolute Scores (Olivoto et al., 2019) for quantifying the stability of g genotypes conducted in e environments using linear mixed-effect models.

The weighted average of absolute scores is computed considering all Interaction Principal Component Axis (IPCA) from the Singular Value Decomposition (SVD) of the matrix of genotype-environment interaction (GEI) effects generated by a linear mixed-effect model, as follows:

$$WAASB_i = \sum_{k=1}^{p} |IPCA_{ik} \times EP_k| / \sum_{k=1}^{p} EP_k$$

where $WAASB_i$ is the weighted average of absolute scores of the *i*th genotype; $IPCA_{ik}$ is the score of the *i*th genotype in the *k*th Interaction Principal Component Axis (IPCA); and EP_k is the explained variance of the *k*th IPCA for k = 1, 2, ..., p, considering p = min(g - 1; e - 1).

The nature of the effects in the model is chosen with the argument random. By default, the experimental design considered in each environment is a randomized complete block design. If block is informed, a resolvable alpha-lattice design (Patterson and Williams, 1976) is implemented. The following six models can be fitted depending on the values of random and block arguments.

• Model 1: block = NULL and random = "gen" (The default option). This model considers a Randomized Complete Block Design in each environment assuming genotype and genotype-environment interaction as random effects. Environments and blocks nested within environments are assumed to fixed factors.

• Model 2: block = NULL and random = "env". This model considers a Randomized Complete Block Design in each environment treating environment, genotype-environment interaction, and blocks nested within environments as random factors. Genotypes are assumed to be fixed factors.

- Model 3: block = NULL and random = "all". This model considers a Randomized Complete Block Design in each environment assuming a random-effect model, i.e., all effects (genotypes, environments, genotype-vs-environment interaction and blocks nested within environments) are assumed to be random factors.
- Model 4: block is not NULL and random = "gen". This model considers an alpha-lattice design in each environment assuming genotype, genotype-environment interaction, and incomplete blocks nested within complete replicates as random to make use of inter-block information (Mohring et al., 2015). Complete replicates nested within environments and environments are assumed to be fixed factors.
- Model 5: block is not NULL and random = "env". This model considers an alpha-lattice design in each environment assuming genotype as fixed. All other sources of variation (environment, genotype-environment interaction, complete replicates nested within environments, and incomplete blocks nested within replicates) are assumed to be random factors.
- **Model 6:** block is not NULL and random = "all". This model considers an alpha-lattice design in each environment assuming all effects, except the intercept, as random factors.

Usage

```
waasb(
   .data,
   env,
   gen,
   rep,
   resp,
   block = NULL,
   by = NULL,
   wresp = NULL,
   random = "gen",
   prob = 0.05,
   ind_anova = FALSE,
   verbose = TRUE,
   ...
)
```

Arguments

.data	The dataset containing the columns related to Environments, Genotypes, replication/block and response variable(s).
env	The name of the column that contains the levels of the environments.
gen	The name of the column that contains the levels of the genotypes.
rep	The name of the column that contains the levels of the replications/blocks.
resp	The response variable(s). To analyze multiple variables in a single procedure a vector of variables may be used. For example $resp = c(var1, var2, var3)$.
block	Defaults to NULL. In this case, a randomized complete block design is considered. If block is informed, then an alpha-lattice design is employed considering

block as random to make use of inter-block information, whereas the complete replicate effect is always taken as fixed, as no inter-replicate information was to be recovered (Mohring et al., 2015).

by One variable (factor) to compute the function by. It is a shortcut to dplyr::group_by().This

is especially useful, for example, when the researcher want to compute the indexes by mega-environments. In this case, an object of class waasb_grouped is returned. mtsi() can then be used to compute the mtsi index within each

mega-environment.

mresp The new maximum value after rescaling the response variable. By default, all

variables in resp are rescaled so that de maximum value is 100 and the minimum value is 0 (i.e., mresp = NULL). It must be a character vector of the same length of resp if rescaling is assumed to be different across variables, e.g., if for the first variable smaller values are better and for the second one, higher values are better, then mresp = c("1,h") must be used. Character value of length 1 will

be recycled with a warning message.

wresp The weight for the response variable(s) for computing the WAASBY index. By

default, all variables in resp have equal weights for mean performance and stability (i.e., wresp = 50). It must be a numeric vector of the same length of resp to assign different weights across variables, e.g., if for the first variable equal weights for mean performance and stability are assumed and for the second one, a higher weight for mean performance (e.g. 65) is assumed, then wresp = c(50,65) must be used. Numeric value of length 1 will be recycled with a

warning message.

random The effects of the model assumed to be random. Defaults to random = "gen".

See **Details** to see the random effects assumed depending on the experimental

design of the trials.

prob The probability for estimating confidence interval for BLUP's prediction.

ind_anova Logical argument set to FALSE. If TRUE an within-environment ANOVA is per-

formed

verbose Logical argument. If verbose = FALSE the code will run silently.

Arguments passed to the function impute_missing_val() for imputation of missing values in the matrix of BLUPs for genotype-environment interaction,

thus all aming the assumptation of the WAACD in dec

thus allowing the computation of the WAASB index.

Value

An object of class waasb with the following items for each variable:

- individual A within-environments ANOVA considering a fixed-effect model.
- fixed Test for fixed effects.
- random Variance components for random effects.
- LRT The Likelihood Ratio Test for the random effects.
- model A tibble with the response variable, the scores of all IPCAs, the estimates of Weighted Average of Absolute Scores, and WAASBY (the index that considers the weights for stability and mean performance in the genotype ranking), and their respective ranks.
- **BLUPgen** The random effects and estimated BLUPS for genotypes (If random = "gen" or random = "all")
- **BLUPenv** The random effects and estimated BLUPS for environments, (If random = "env" or random = "all").

- **BLUPint** The random effects and estimated BLUPS of all genotypes in all environments.
- **PCA** The results of Principal Component Analysis with the eigenvalues and explained variance of the matrix of genotype-environment effects estimated by the linear fixed-effect model.
- MeansGxE The phenotypic means of genotypes in the environments.
- **Details** A list summarizing the results. The following information are shown: Nenv, the number of environments in the analysis; Ngen the number of genotypes in the analysis; mresp The value attributed to the highest value of the response variable after rescaling it; wresp The weight of the response variable for estimating the WAASBY index. Mean the grand mean; SE the standard error of the mean; SD the standard deviation. CV the coefficient of variation of the phenotypic means, estimating WAASB, Min the minimum value observed (returning the genotype and environment), Max the maximum value observed (returning the genotype and environment); MinENV the environment with the lower mean, MaxENV the environment with the larger mean observed, MinGEN the genotype with the lower mean, MaxGEN the genotype with the larger.
- **ESTIMATES** A tibble with the genetic parameters (if random = "gen" or random = "all") with the following columns: Phenotypic variance the phenotypic variance; Heritability the broad-sense heritability; GEr2 the coefficient of determination of the interaction effects; h2mg the heritability on the mean basis; Accuracy the selective accuracy; rge the genotype-environment correlation; CVg the genotypic coefficient of variation; CVr the residual coefficient of variation; CV ratio the ratio between genotypic and residual coefficient of variation.
- residuals The residuals of the model.
- formula The formula used to fit the model.

Author(s)

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References

Olivoto, T., A.D.C. L\'ucio, J.A.G. da silva, V.S. Marchioro, V.Q. de Souza, and E. Jost. 2019. Mean performance and stability in multi-environment trials I: Combining features of AMMI and BLUP techniques. Agron. J. 111:2949-2960. doi: 10.2134/agronj2019.03.0220

Mohring, J., E. Williams, and H.-P. Piepho. 2015. Inter-block information: to recover or not to recover it? TAG. Theor. Appl. Genet. 128:1541-54. doi: 10.1007/s0012201525300

Patterson, H.D., and E.R. Williams. 1976. A new class of resolvable incomplete block designs. Biometrika 63:83-92.

See Also

```
mtsi() waas() get_model_data() plot_scores()
```

```
rep = REP,
           resp = everything())
# Distribution of random effects (first variable)
plot(model, type = "re")
# Genetic parameters
get_model_data(model, "genpar")
# Example 2: Analyzing variables that starts with "N" #
# assuming environment as random effects with higher weight for #
# response variable (65) for the three traits.#
model2 <- waasb(data_ge2,</pre>
            env = ENV,
            gen = GEN,
            rep = REP,
            random = "env",
            resp = starts_with("N"),
            wresp = 65)
# Get the index WAASBY
get_model_data(model2, what = "WAASBY")
# Plot the scores (response x WAASB)
plot_scores(model2, type = 3)
# Example 3: Analyzing GY and HM assuming a random-effect model.#
# Smaller values for HM and higher values for GY are better. #
# To estimate WAASBY, higher weight for the GY (60%) and lower #
# weight for HM (40%) are considered for mean performance.
#----#
model3 <- waasb(data_ge,</pre>
            env = ENV,
            gen = GEN,
            rep = REP,
            resp = c(GY, HM),
            random = "all",
            mresp = c("h, 1"),
            wresp = c(60, 40))
# Get Likelihood-ratio test
get_model_data(model3, "lrt")
# Get the random effects
get_model_data(model3, what = "ranef")
```

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waas_means

Weighted Average of Absolute Scores

Description

[Stable]

Compute the Weighted Average of Absolute Scores (Olivoto et al., 2019) based on means for genotype-environment data as follows:

$$WAAS_i = \sum_{k=1}^{p} |IPCA_{ik} \times EP_k| / \sum_{k=1}^{p} EP_k$$

where $WAAS_i$ is the weighted average of absolute scores of the *i*th genotype; PCA_{ik} is the score of the *i*th genotype in the *k*th IPCA; and EP_k is the explained variance of the *k*th IPCA for k = 1,2,...,p, where p is the number of IPCAs that explain at least an amount of the genotype-interaction variance declared in the argument min_expl_var.

Usage

```
waas_means(
   .data,
   env,
   gen,
   resp,
   mresp = NULL,
   wresp = NULL,
   win_expl_var = 85,
   verbose = TRUE,
   ...
)
```

Arguments

cation/block and response variable(s).

env The name of the column that contains the levels of the environments.

gen The name of the column that contains the levels of the genotypes.

resp The response variable(s). To analyze multiple variables in a single procedure

a vector of variables may be used. For example resp = c(var1, var2, var3).

Select helpers are also allowed.

mresp The new maximum value after rescaling the response variable. By default, all

variables in resp are rescaled so that de maximum value is 100 and the minimum value is 0 (i.e., mresp = NULL). It must be a character vector of the same length of resp if rescaling is assumed to be different across variables, e.g., if for the first variable smaller values are better and for the second one, higher values are better, then mresp = c("1,h") must be used. Character value of length 1 will

be recycled with a warning message.

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wresp The weight for the response variable(s) for computing the WAASBY index.

Must be a numeric vector of the same length of resp. Defaults to 50, i.e., equal

weights for stability and mean performance.

min_expl_var The minimum explained variance. Defaults to 85. Interaction Principal Compo-

ment Axis are iteractively retained up to the explained variance (eigenvalues in the singular value decomposition of the matrix with the interaction effects) be greather than or equal to min_expl_var. For example, if the explained variance (in percentage) in seven possible IPCAs are 56, 21, 9, 6, 4, 3, 1, resulting in a cumulative proportion of 56, 77, 86, 92, 96, 99, 100, then p = 3, i.e., three

IPCAs will be used to compute the index WAAS.

verbose Logical argument. If verbose = FALSE the code is run silently.

... Arguments passed to the function impute_missing_val() for imputation of

missing values in case of unbalanced data.

Value

An object of class waas_means with the following items for each variable:

- model A data frame with the response variable, the scores of all Principal Components, the
 estimates of Weighted Average of Absolute Scores, and WAASY (the index that consider the
 weights for stability and productivity in the genotype ranking.
- **ge_means** A tbl_df containing the genotype-environment means.
- **ge_eff** A gxe matrix containing the genotype-environment effects.
- **eigenvalues** The eigenvalues from the singular value decomposition of the matrix withe the genotype-environment interaction effects.
- **proportion** The proportion of the variance explained by each IPCA.
- cum_proportion The cumulative proportion of the variance explained.

Author(s)

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References

Olivoto, T., A.D.C. L\'ucio, J.A.G. da silva, V.S. Marchioro, V.Q. de Souza, and E. Jost. 2019a. Mean performance and stability in multi-environment trials I: Combining features of AMMI and BLUP techniques. Agron. J. 111:2949-2960. doi: 10.2134/agronj2019.03.0220

See Also

```
waas() waasb()
```

266 wsmp

wsmp

Weighting between stability and mean performance

Description

[Stable]

This function computes the WAASY or WAASBY indexes (Olivoto et al., 2019) considering different scenarios of weights for stability and mean performance.

After fitting a model with the functions waas() or waasb() it is possible to compute the superiority indexes WAASY or WAASBY in different scenarios of weights for stability and mean performance. The number of scenarios is defined by the arguments increment. By default, twenty-one different scenarios are computed. In this case, the the superiority index is computed considering the following weights: stability (waasb or waas) = 100; mean performance = 0. In other words, only stability is considered for genotype ranking. In the next iteration, the weights becomes 95/5 (since increment = 5). In the third scenario, the weights become 90/10, and so on up to these weights become 0/100. In the last iteration, the genotype ranking for WAASY or WAASBY matches perfectly with the ranks of the response variable.

Usage

```
wsmp(
  model,
  mresp = 100,
  increment = 5,
  saveWAASY = 50,
  prob = 0.05,
  progbar = TRUE
)
```

Arguments

model

Should be an object of class waas or waasb.

mresp

A numeric value that will be the new maximum value after rescaling. By default, the variable in resp is rescaled so that the original maximum and minimum values are 100 and 0, respectively. Let us consider that for a specific trait, say, lodging incidence, lower values are better. In this case, you should use mresp = 0 to rescale the response variable so that the lowest values will become 100 and the highest values 0.

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increment	The increment in the weight ratio for stability and mean performance. Se the Details section for more information.
saveWAASY	Automatically save the WAASY values when the weight for stability is saveWAASY. Default is 50. Please, note that saveWAASY
prob	The p-value for considering an interaction principal component axis significant. must be multiple of increment. If this assumption is not valid, an error will be occur.
progbar	A logical argument to define if a progress bar is shown. Default is TRUE.

Value

An object of class wsmp with the following items for each variable:

- scenarios A list with the model for all computed scenarios.
- **WAASY** The values of the WAASY estimated when the weight for the stability in the loop match with argument saveWAASY.
- hetdata, hetcomb The data used to produce the heatmaps.
- Ranks All the values of WAASY estimated in the different scenarios of WAAS/GY weighting ratio.

Author(s)

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References

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