# Package 'pepa'

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Type Package

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| Title Package for the Execution of Pre Cooked Analysis   |
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| <b>Description</b> This packages creates automatic reports for different types of statistical methodologies. |
| <b>Depends</b> R (>= 3.0.0), st4gi   |
| Imports rmarkdown  |
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| NeedsCompilation no  |
| R topics documented:   |
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pty

Pepa tells you

# **Description**

Explain an R object in plain English if she knows about it

# Usage

```
pty(x, author = "International Potato Center")
```

# Arguments

x An R object. author Author.

#### **Details**

It uses a set of templates to explain R objects in plain English. It aims to produce automatic reports for some standard statistical procedures, most of them included in the st4gi package.

#### Value

It returns an explanation about the selected R object.

# Author(s)

Raul Eyzaguirre.

# **Examples**

```
# Pepa tells you something about a data frame:
pty(pjpz09)
```

pty.aovmet

Pepa tells you about a MET with a RCBD

# **Description**

Explain a fitted model for a multi environment trial (MET) with a RCBD in each environment in plain English.

# Usage

```
pty.aovmet(trait, geno, env, rep, data, maxp = 0.1,
  author = "International Potato Center")
```

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# **Arguments**

| trait  | The trait to analize.   |
|--------|---|
| geno   | The genotypes.  |
| env    | The environments.   |
| rep    | The replications.   |
| data   | The name of the data frame containing the data.                           |
| maxp   | Maximum allowed proportion of missing values to estimate, default is 10%. |
| author | Author.   |

#### **Details**

It fits a linear model for a MET with a RCBD and explains the results. If data is unbalanced, missing values are estimated up to an specified maximum proportion, 10% by default. Genotypes and environments are considered as fixed factors while the blocks are considered as random and nested into the environments.

# Value

It returns an explanation about the MET with a RCBD fitted model.

# Author(s)

Raul Eyzaguirre.

# **Examples**

```
pty.aovmet("y", "geno", "env", "rep", met8x12)
```

pty.elston

Pepa tells you about the Elston index

# **Description**

Explain the results of the Elston index in plain English.

# Usage

```
pty.elston(traits, geno, env = NULL, rep = NULL, data, means = "single",
  model = "gxe", lb = 1, author = "International Potato Center")
```

#### **Arguments**

| traits | List of traits.                                     |
|--------|---|
| geno   | The genotypes.                                      |
| env    | The environments.                                   |
| rep    | The replications. Must be defined if model = "gxe". |
| data   | The name of the data frame containing the data.     |

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| means  | The genotypic means to compute the index, "single" or "fitted". The default is "single". See details for more information.   |
|--------|--|
| model  | Type of model to fit means if means = "fitted", "gxe" for a model with gxe interaction or "g+e" for a model without interaction. The default is "gxe". See details for more information. |
| lb     | Lower bound. 1 for $k = min(x)$ and 2 for $k = (n \times min(x) - max(x))/(n-1)$   |
| author | Author.  |

#### **Details**

The Elston index is a weight free index.

If means = "fitted" and model = "gxe" then the arguments env and rep must be specified. If means = "fitted" and model = "g+e" then only the argument env must be specified. If means = "single" and env and rep are specified, then single arithmetic means are computed over the replications for each genotype at each environment and then for each genotype over environments. In any other case single arithmetic means are computed over all the observations for each genotype.

#### Value

It returns an explanation about the Elston index.

#### Author(s)

Raul Eyzaguirre.

# **Examples**

```
pty.elston(c("rytha", "bc", "dm", "star", "nocr"), "geno", data = spg)
```

pty.pesekbaker

Pepa tells you about the Pesek-Baker index

# **Description**

Explain the results of the Pesek-Baker index in plain English.

## Usage

```
pty.pesekbaker(traits, geno, env, rep = NULL, data, means = "single",
  model = "gxe", dgg = NULL, units = "sdu", sf = 0.1,
  author = "International Potato Center")
```

#### **Arguments**

| traits | List of traits.                                     |
|--------|---|
| geno   | The genotypes.                                      |
| env    | The environments.                                   |
| rep    | The replications. Must be defined if model = "gxe". |
| data   | The name of the data frame containing the data.     |

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| means  | The genotypic means to compute the index, "single" or "fitted". The default is "single". See details for more information.                              |
|--------|---|
| model  | Type of model, "gxe" for a model with gxe interaction or "g+e" for a model without interaction. The default is "gxe". See details for more information. |
| dgg    | Desired genetic gains. The default is one standard deviation for each trait.  |
| units  | Units for dgg, "actual" or "sdu". See details for more information.   |
| sf     | Selected fraction. The default is 0.1.  |
| author | Author.   |

**Details** 

The Pesek-Baker is an index where relative economic weights have been replaced by desired gains.

By default a model with components for genotypes, environments, genotypes by environments interaction and replications nested into environments is fitted (model = "gxe"). If model = "g+e" then a model with components for genotypes and environments is fitted, and in this case the gxe variance includes the gxe plus the error variance. Response to selection is only computed when model = "gxe".

If means = "fitted" then the model specified in model is used to fit the means of the genotypes. Otherwise single arithmetic means are computed over the replications for each genotype at each environment and then for each genotype over environments.

If dgg is not specified, the standard deviations of the traits are used. It means that the desired genetic gains are equal to one standard deviation for each trait. dgg can be specified in actual units (units = "actual") or in standard deviations (units = "sdu"), defaults to "sdu". For example, if you have a trait which is expressed in kilograms and with a standard deviation of 5 kilograms, typing dgg = 2 means a desired genetic gain of 2 standard deviations that corresponds to 10 kilograms. If you type dgg = 2 and units = "actual" then this means a desired genetic gain of 2 kilograms. If dgg = NULL then the desired genetic gain will be one standard deviation, no matter if units is set as "actual" or "sdu".

#### Value

It returns an explanation about the Pesek-Baker index.

# Author(s)

Raul Eyzaguirre.

#### **Examples**

```
pty.pesekbaker(c("rytha", "bc", "dm", "star", "nocr"), "geno", "loc", "rep", spg)
```

pty.rcbd

Pepa tells you about RCBD

# **Description**

Explain a RCBD fitted model in plain English

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#### Usage

```
pty.rcbd(trait, treat, rep, data, maxp = 0.1,
  author = "International Potato Center")
```

# **Arguments**

trait The trait to analize.
treat The treatments.
rep The replications.

data The name of the data frame.

maxp Maximum allowed proportion of missing values to estimate, default is 10%.

author Author.

#### **Details**

It fits a linear model for a RCBD and explains the results. It also checks the assumptions.

#### Value

It returns an explanation about the RCBD fitted model.

# Author(s)

Raul Eyzaguirre.

# **Examples**

```
pty.rcbd("trw", "geno", "rep", pjpz09)
```

pty.spconsis

Pepa tells you about the consistency of your data

# Description

It checks your data for inconsistencies.

# Usage

```
pty.spconsis(data, plot.size, f = 3, author = "International Potato Center")
```

# **Arguments**

data The name of the data frame.

plot.size Plot size in square meters.

f Factor for extreme values detection. See details.

author Author.

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#### **Details**

The data frame must use the labels (lower or upper case) specified in the function spconsis of package st4gi. Type ?spconsis to see the list. Extreme values are detected using the interquartile range. The rule is to detect any value out of the interval  $[Q_1 - f \times IQR; Q_3 + f \times IQR]$ . By default f = 3.

# Value

It returns a list of all rows with some kind of inconsistency and all rows with outliers.

#### Author(s)

Raul Eyzaguirre.

## **Examples**

```
pty.spconsis(pjpz09, 4.5)
```

repo.met

Authomatic report for a MET with a RCBD

#### **Description**

Produces an authomatic report for a selected trait in a multi environment trial (MET) with a RCBD in each environment.

### Usage

```
repo.met(trait, geno, env, rep, data, maxp = 0.1,
  author = "International Potato Center")
```

# **Arguments**

| trait  | The trait to be evaluated.  |
|--------|---|
| geno   | The genotypes.  |
| env    | The environments.   |
| rep    | The replications.   |
| data   | The name of the data frame containing the data.                           |
| maxp   | Maximum allowed proportion of missing values to estimate, default is 10%. |
| author | Author.   |

#### **Details**

It fits a linear model for a MET with a RCBD for the selected trait. If data is unbalanced, missing values are estimated up to an specified maximum proportion, 10% by default. Genotypes and environments are considered as fixed factors while the blocks are considered as random and nested into the environments.

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# Value

It returns an automatic report about the MET with a RCBD fitted model.

# Author(s)

Raul Eyzaguirre.

# Examples

```
repo.met("rytha", "geno", "env", "rep", megaclones)
```

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