# Package 'Phenotype'

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Type Package

Title A Tool for Phenotypic Data Processing

Version 0.1.0

Maintainer Peng Zhao <pengzhao@nwafu.edu.cn>

Description Large-scale phenotypic data processing is essential in research. Researchers need to eliminate outliers from the data in order to obtain true and reliable results. Best linear unbiased prediction (BLUP) is a standard method for estimating random effects of a mixed model. This method can be used to process phenotypic data under different conditions and is widely used in animal and plant breeding. The 'Phenotype' can remove outliers from phenotypic data and performs the best linear unbiased prediction (BLUP), help researchers quickly complete phenotypic data analysis. H.P.Piepho. (2008) <doi:10.1007/s10681-007-9449-8>.

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**Encoding** UTF-8

LazyData true

URL https://github.com/biozhp/Phenotype

BugReports https://github.com/biozhp/Phenotype/issues

Depends R (>= 3.5.0) Imports lme4, tidyr RoxygenNote 7.1.1 NeedsCompilation no Author Peng Zhao [aut, cre] Repository CRAN Date/Publication 2020-08-06 14:40:19 UTC

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blup

blup

# Description

Performs the Best Linear Unbiased Prediction (BLUP)

#### Usage

```
blup(
  х,
  sample = NULL,
  year = NULL,
  loc = NULL,
  rep = NULL,
  phe = NULL,
  fold = 1.5
```

#### Arguments

)

х	Input phenotype data file.
sample	The column name of the sample name in phenotypic data. (Default: NULL)
year	The column name of the year in phenotypic data. (Default: NULL)
loc	The column name of the location in phenotypic data. (Default: NULL)
rep	The column name of the replication in phenotypic data. (Default: NULL)
phe	The column name of the phenotypic value in data. (Default: NULL)
fold	Fold before inter-quartile range. (Default: 1.5)

# Value

Estimate BLUPs for a phenotypic data with outliers removed on a per sample basis.

# Author(s)

Peng Zhao <pengzhao@nwafu.edu.cn>

# Examples

```
data("wheatds")
blup_out <- blup(wheatds, sample = "Line", loc = "Env", rep = "Rep", phe = "DS")</pre>
```

blup

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histplot

histplot

# Description

Histogram drawing

## Usage

```
histplot(
  х,
  color = "#99d6e1",
  rug_color = "#f79999",
  freq = FALSE,
  1wd = 2,
  rug_{lwd} = 1,
  main = "",
xlab = "",
 ylab = "",
  cex.main = 1.5,
  cex.lab = 1.5,
  cex.axis = 1.5,
  breaks = "Sturges",
  ylim = NULL,
  xpos = 0.03,
  ypos = 0,
  cex.text = 1.2
)
```

# Arguments

Х	Input phenotype data.
color	The color of histogram.
rug_color	The color of rug under the histogram.
freq	If FALSE, the histogram graphic is a representation of frequencies; if TRUE, the histogram graphic is a representation of probability densitie. (Default: FALSE)
lwd	The line width of histogram. (Default: 2)
rug_lwd	The line width of rug under the histogram. (Default: 1)
main	The title of plot.
xlab	The X axis labels.
ylab	The Y axis labels
cex.main	The magnification to be used for title. (Default: 1.5)
cex.lab	The magnification to be used for axis labels. (Default: 1.5)
cex.axis	The magnification to be used for axis annotation. (Default: 1.5)

outlier

breaks	The number of bars in the histogram.
ylim	Y axis ranges.
xpos	The horizontal position of the pvalue label. (Default: 0.03)
ypos	The vertical position of the pvalue label. (Default: 0)
cex.text	The magnification to be used for pvalue labels. (Default: 1.2)

#### Value

Histogram and p-value of Shapiro-Wilk Normality Test.

#### Author(s)

Peng Zhao <pengzhao@nwafu.edu.cn>

#### Examples

```
data("wheatds")
inlier <- outlier(wheatds, sample = "Line", loc = "Env", rep = "Rep", phe = "DS", mode = "blup")
stat_out <- stat(x = inlier, sample = "Sample", phe = "inlier")
histplot(x = stat_out$mean)</pre>
```

outlier

outlier

#### Description

Remove outliers from phenotypic data

#### Usage

```
outlier(
    x,
    sample = NULL,
    year = NULL,
    loc = NULL,
    rep = NULL,
    phe = NULL,
    fold = 1.5,
    mode = "normal"
)
```

stat

# Arguments

х	Input phenotype data file.
sample	The column name of the sample name in phenotypic data. (Default: NULL)
year	The column name of the year in phenotypic data. (Default: NULL)
loc	The column name of the location in phenotypic data. (Default: NULL)
rep	The column name of the replication in phenotypic data. (Default: NULL)
phe	The column name of the phenotypic value in data. (Default: NULL)
fold	Fold before inter-quartile range. (Default: 1.5)
mode	Type of input phenotypic data. "normal" means normal data, "blup" means data containing year/location/repetition. (Default: "normal")

# Value

phenotypic data with outliers removed.

#### Author(s)

Peng Zhao <pengzhao@nwafu.edu.cn>

# Examples

```
data("wheatds")
inlier <- outlier(wheatds, sample = "Line", loc = "Env", rep = "Rep", phe = "DS", mode = "blup")</pre>
```

# Description

Calculate statistical indicators of phenotypic data

# Usage

stat(x, sample = NULL, phe = NULL)

#### Arguments

Х	Input phenotype data file.
sample	The column name of the sample name in phenotypic data. (Default: NULL)
phe	The column name of the phenotypic value in data. (Default: NULL)

## Value

Mean, median, standard deviation, standard error of phenotypic data for each sample.

## Author(s)

Peng Zhao <pengzhao@nwafu.edu.cn>

#### Examples

```
data("wheatds")
inlier <- outlier(wheatds, sample = "Line", loc = "Env", rep = "Rep", phe = "DS", mode = "blup")
stat_out <- stat(x = inlier, sample = "Sample", phe = "inlier")</pre>
```

wheatds	Stripe rust disease severity (leaf areas infected, DS) of the wheat RIL
	population

#### Description

Stripe rust disease severity (leaf areas infected, DS) of the wheat RIL population in Yangling, Tianshui, Jiangyou.

# Usage

data("wheatds")

#### Format

A data frame containing samples, environments, repetitions, and disease severity of the wheat RIL population.

## Examples

data("wheatds")

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