Package 'gwaRs'

October 13, 2022

Type Package

Title Manhattan, Q-Q, and PCA Plots using 'ggplot2'

Version 0.3.0

Description Generate Manhattan, Q-Q, and PCA plots from GWAS and PCA results using 'ggplot2'.

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Encoding UTF-8

LazyData true

Imports RColorBrewer (>= 1.1.2), ggplot2, ggrepel, dplyr, tidyr, data.table, scales, stats, grDevices

RoxygenNote 7.1.1

Depends R (>= 2.10)

Suggests knitr, rmarkdown

VignetteBuilder knitr

URL https://github.com/LindoNkambule/gwaRs

BugReports https://github.com/LindoNkambule/gwaRs/issues

NeedsCompilation no

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Repository CRAN

Date/Publication 2021-04-19 17:30:06 UTC

R topics documented:

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```
gwaRs
```

Create Manhattan, Q-Q, and PCA plots for GWAS data.

Description

A package for creating Manhattan, Q-Q, and PCA plots for GWAS data.

Author(s)

Lindokuhle Nkambule <lindonkambule116@gmail.com>

See Also

Useful links:

- https://github.com/LindoNkambule/gwaRs
- Report bugs at https://github.com/LindoNkambule/gwaRs/issues

gwasData

GWAS results

Description

Example GWAS data.

highlightSNPS Highlight SNPs

Description

A couple of SNPs to be annotated

karyotype_plot Karyotype Plot

Description

Creates a Karyotype plot

Usage

```
karyotype_plot(
   data,
   density.col = c("darkgreen", "yellow", "red"),
   window.size = 1e+06,
   title = NULL
)
```

Arguments

data	A data.frame with "CHR" and "BP"columns.
density.col	A character vector with colors to use for gradients.
window.size	A double precision numeric value indicating the window size.
title	A string denoting the title to use for the plot. Default is 'Manhattan Plot'

Details

Creates a SNP Karyotype or Density plot from an R dataframe with "CHR" and "BP" columns.

Value

A SNP Karyotype plot.

Author(s)

Lindokuhle Nkambule

Examples

karyotype_plot(gwasData)

man_plot

Description

Creates a Manhattan plot

Usage

```
man_plot(
    data,
    chromCol = c("gray44", "black"),
    genomewideline = -log10(5e-08),
    suggestiveline = -log10(1e-05),
    chromosome = "ALL",
    annotatePval = FALSE,
    annotateSNP = NULL,
    annotateCol = "red",
    highlight = NULL,
    highlightCol = "green3",
    title = NULL
)
```

Arguments

PLINK assoc output, tab-delimited, or a data.frame with "SNP", "CHR", "BP", and "P" columns.
A character vector indicating which colors to alternate for the chromosomes.
Where to draw the "genome-wide significant" line. Default -log10(5e-8). Set to FALSE or F to disable
Where to draw the "suggestive" line. Default -log10(1e-5). Set to FALSE or F to disable.
An integer indicating which chromosome to plot. Default is "ALL".
If set, SNPs with p-value less than or equal to this p-value will be annotated on the plot.
A character vector of SNPs in your dataset to annotate. If some of the SNPs are not in your dataset, gwaRs will throw a warning message.
A string denoting the color to use for the annotations.
A character vector of SNPs in the dataset to highlight. If some of the SNPs are not in your dataset, gwaRs will throw a warning message. Default is NULL.
A string denoting the color to use to highlight the SNPs.
A string denoting the title to use for the plot. Default is 'Manhattan Plot'

Details

Creates a Manhattan plot from PLINK assoc output (or any tab-delimited file or data frame with "SNP", "CHR", "BP", and "P" columns).

Value

A Manhattan plot.

Author(s)

Lindokuhle Nkambule

Examples

man_plot(gwasData)

mirrored_man_plot Mirrored Manhattan Plot

Description

Creates a Mirrored Manhattan Plot for two traits

Usage

```
mirrored_man_plot(
  data,
  trait1 = NULL,
  trait2 = NULL,
  trait1_chromCols = c("gray66", "grey36"),
  trait2_chromCols = c("steelblue1", "steelblue4"),
  xlab = "Genomic Position (chromosome)",
  title = "Manhattan Plot",
  annotate_trait1_pval = FALSE,
  annotate_trait1_color = "red",
  annotate_trait2_pval = FALSE,
  annotate_trait2_color = "red",
  annotateSNP = NULL,
  annotateSNPcolor = "red",
  highlight = NULL,
  highlightcolor = "green3",
  genomewideline_trait1 = NULL,
  genomewideline_trait2 = NULL,
  genomewideline_type = "dashed",
  genomewideline_color = "red",
  suggestiveline_trait1 = NULL,
```

```
suggestiveline_trait2 = NULL,
suggestiveline_type = "dashed",
suggestiveline_color = "blue"
)
```

Arguments

data	A tab-delimited or data frame with the compulsory columns: "CHR", "SNP", "BP", "P", "Trait".							
trait1	A character string of the trait1 as it appears in the input data.							
trait2	A character string of the trait2 as it appears in the input data.							
<pre>trait1_chromCol</pre>	ls							
	A character vector indicating which colors to alternate for trait1 chromosomes.							
trait2_chromCol								
	A character vector indicating which colors to alternate for trait2 chromosomes.							
xlab	A character string to be used as the x-axis label.							
title	A character string to be used as the plot title							
annotate_trait1	I_pval							
	If set, trait1 SNPs with p-value less than or equal to this p-value will be annotated on the plot.							
annotate_trait1	I_color							
	A character string indicating the color to be used for annotating trait1 SNPs by p-value							
annotate_trait2_pval								
	If set, trait2 SNPs with p-value less than or equal to this p-value will be annotated on the plot.							
annotate_trait2	2_color							
	A character string indicating the color to be used for annotating trait2 SNPs by p-value							
annotateSNP	A character vector of SNPs in your dataset to annotate. If some of the SNPs are not in your dataset, gwaRs will throw a warning message.							
annotateSNPcolo	or							
	A character string denoting the color to use for the annotations.							
highlight	A character vector of SNPs in the dataset to highlight. If some of the SNPs are not in your dataset, gwaRs will throw a warning message. Default is NULL.							
highlightcolor	A character string denoting the color to use to highlight the SNPs.							
genomewideline_	_trait1							
-	Where to draw the "genome-wide significant" line for trait1							
genomewideline_	_trait2							
	Where to draw the "genome-wide significant" line for trait2							
genomewideline_	_type							
	A character string denoting the type of line to be used for the "genome-wide significant" line. This is the same for both traits. Default is dashed.							

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pcaData

genomewideline_color

A character string denoting the color to be used for the "genome-wide significant" line. This is the same for both traits. Default is red.

suggestiveline_trait1

Where to draw the "suggestive" line for trait1.

suggestiveline_trait2

Where to draw the "suggestive" line for trait2.

suggestiveline_type

A character string denoting the type of line to be used for the "suggestive" line. This is the same for both traits. Default is dashed

suggestiveline_color

A character string denoting the color to be used for the "suggestive" line. This is the same for both traits. Default is blue.

Details

Create a Mirrored Manhattan Plot from a tab-delimited file or data frame with the compulsory columns: "CHR", "SNP", "BP", "P", "Trait".

Value

A Mirrored Manhattan plot for two traits.

Author(s)

Lindokuhle Nkambule

Examples

```
## Not run:
mirrored_man_plot(inputData)
```

End(Not run)

pcaData

PCA results

Description

Example PCA data.

pca_plot

PCA Plot

Description

Creates a PCA plot

Usage

```
pca_plot(
    data,
    xComponent = "PC1",
    yComponent = "PC2",
    legendPos = "right",
    soft = "PLINK",
    colPalette = "Accent",
    title = NULL
)
```

Arguments

data	PLINK pca or EIGENSTRAT smartpca output, (or any tab-delimited file or data.frame with the same format as PLINK pca or EIGENSTRAT smartpca output)
xComponent	A character vector indicating the principal component value to use for the x-axis. Default is "PC1"
yComponent	A character vector indicating the principal component value to use for the x-axis. Default is "PC2"
legendPos	A character vector indicating the legend position. Default is "right".
soft	A character vector indicating the software output format. Default is "PLINK". If you have a tab-delimited file or data.frame with the same format as EIGEN-STRAT output, use "EIGENSTRAT"
colPalette	A character vector indicating the color palette to use. Default is "Accent".
title	A string denoting the title to use for the plot. Default is 'PCA Plot'

Details

Creates a PCA plot from PLINK pca output, EIGENSTRAT smartpca, (or any tab-delimited file or data.frame with the same format as PLINK pca or EIGENSTRAT smartpca output).

Value

A PCA plot.

Author(s)

Lindokuhle Nkambule

qq_plot

Examples

pca_plot(pcaData)

qq_plot

QQ Plot

Description

Creates a Q-Q plot

Usage

```
qq_plot(
   data,
   point_col = "black",
   diag_col = "red",
   diag_line = "solid",
   title = NULL
)
```

Arguments

data	PLINK assoc output, tab-delimited, or a data.frame with "P" column.
point_col	A character vector indicating the color to use for the SNP p-values. Default is "black".
diag_col	A character vector indicating the color to use for the diagonal line. Default is "red".
diag_line	A character vector indicating the line type to use for the diagonal line. Default is "solid".
title	A string denoting the title to use for the plot. Default is 'Q-Q Plot'

Details

Creates a Q-Q plot from PLINK assoc output (or any tab-delimited file or data frame with "P" column).

Value

A Q-Q plot.

Author(s)

Lindokuhle Nkambule

qq_plot

Examples

qq_plot(gwasData)

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