# Package 'bugwas'

March 15, 2016				
Title What the Package Does (one line, title case) Version 0.0.0.9000				
<b>Depends</b> R (>= 3.2.1)				
License What license is it under?				
LazyData true				
Imports ape,				
phangorn				
RoxygenNote 5.0.1.9000				
R topics documented:				
all_plots bayesianWaldTestPCsBarplot genVarPlots lin_loc logregVsLMM pcLoadingPlot plotIndividualBy2PCs snpManhattanPlot snpPCManhattanPlot testGenomeWidePCs testGenomeWidePCs testGenomeWidePCsAll trueAndPredPhenoOnTreePlot  Index	1			
all_plots Generates all plots.				
Description This function generates all the plots Usage				
all plots(hiallelic = NUUL triallelic = NUUL gapVars = NUUL				

treeInfo = NULL, config = NULL)

#### **Arguments**

biallelic	A list called 'biallelic' created from the lin_loc function
triallelic	A list called 'triallelic' created from the lin_loc function
genVars	A list called 'genVars' created from the lin_loc function
treeInfo	A list called 'treeInfo' created from the lin_loc function
config	A list called 'config' created from the lin_loc function

#### **Examples**

```
data <- lin_loc(gen = gen, pheno = pheno, phylo = phylo,
  prefix = prefix, gem.path = gem.path)
all_plots(biallelic = data$biallelic, triallelic = data$triallelic,
genVars = data$genVars, treeInfo = data$treeInfo, config = data$config)</pre>
```

bayesianWaldTestPCsBarplot

Barplot of Bayesian Wald Test on principal components

# **Description**

This function generates the barplot of Bayesian Wald Test on principal components.

# Usage

```
bayesianWaldTestPCsBarplot(config, biallelic, treeInfo, colourPalette = NULL,
    p.genomewidepc = NULL)
```

#### **Arguments**

config A list called 'config' created from the lin\_loc function. It is a required input.

biallelic A list called 'biallelic' created from the lin\_loc function. It is a required input.

treeInfo A list called 'treeInfo' created from the lin\_loc function. It is a required input.

colourPalette A vector of colours colour the significant principal components identified by the Bayesian Wald test (see testGenomeWidePCs). If this is NULL then colours are chosen from a default colour palette.

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p.genomewidepc A matrix of the significant principal component and their correlation with lineages. This is Bayesian Wald test results produced by the function testGenomeWide-PCs. If this is NULL then testGenomeWidePCs is called to generate the required

test results.

```
data <- lin_loc(gen = gen, pheno = pheno, phylo = phylo,
    prefix = prefix, gem.path = gem.path)
testGenomeWidePCs(config = data$config, biallelic = data$biallelic)</pre>
```

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genVarPlots	Plots for GWAS on general variants

# **Description**

This function generates the various Manhattan plots for general variants.

# Usage

```
genVarPlots(genVars, biallelic, config, colourPalette = NULL)
```

# **Arguments**

genVars	A list called 'genVars' created from the lin_loc function. It is a required input.
biallelic	A list called 'biallelic' created from the lin_loc function. It is a required input.
config	A list called 'config' created from the lin_loc function. It is a required input.
colourPalette	A vector of colours colour the significant principal components identified by the Bayesian Wald test (see testGenomeWidePCs). If this is NULL then colours are chosen from a default colour palette.

# **Examples**

```
data <- lin_loc(gen = gen, pheno = pheno, phylo = phylo,
    prefix = prefix, gem.path = gem.path)
genVarPlots(genVars = data$genVars, biallelic = data$biallelic, config = data$config)</pre>
```

lin\_loc

Lineage and locus tests for bacterial GWAS

# **Description**

This function tests for locus effects using GEMMA and lineage effects using a bayesian wald test for haploid data

# Usage

```
lin_loc(gen = NULL, pheno = NULL, phylo = NULL, prefix = NULL,
  gem.path = NULL, pcs = NULL, lmm.bi = NULL, lmm.tri.tetra = NULL,
  logreg.bi = NULL, logreg.tri.tetra = NULL, var.matrix = NULL,
  logreg.var = NULL, lmm.var = NULL, cutOffCor = 0, run.lmm = TRUE,
  maf = 0, relmatrix = NULL, lognull = NULL, lambda = NULL,
  output.dir = getwd(), creatingAllPlots = TRUE,
  allBranchAndPCCor = FALSE, runTriTetrallelic = TRUE)
```

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#### **Arguments**

gen A file name specified by either a variable of mode character, or a double-quoted

string, containing imputed haploid SNP data. Rows are SNPs, and columns are samples, with the first column being SNP positions. Column headers must contain 'ps' for the SNP positions with the others being the sample names. This must contain biallelic SNPs, but can also contain tri and tetra-allelic SNPs. Re-

quired argument.

pheno A file name specified by either a variable of mode character, or a double-quoted

string, containing a column of sample names with header 'ID' and a column of the binary phenotype (coded by 0s and 1s) with column header 'pheno'. Re-

quired argument.

phylo A file name specified by either a variable of mode character, or a double-quoted

string, containing a phylogeny of the samples, with the same names matching

with arguments gen and pheno. Required argument.

prefix Output file prefix. Required argument.

gem.path A file path specified by either a variable of mode character, or a double-quoted

string. gem.path is the file path to the software GEMMA (version >= ?). Re-

quired argument.

pcs A file name specified by either a variable of mode character, or a double-quoted

string, containing the principle components of the data. Column names should

be 'PC1' to 'PCn' and row names should be the sample names.

1mm.bi A file name specified by either a variable of mode character, or a double-quoted

string, containing GEMMA results (ending '.assoc.txt') for the biallelic SNPs in

argument 'gen'.

1mm.tri.tetra A file name specified by either a variable of mode character, or a double-quoted

string, containing GEMMA results for the tri and tetra allelic SNPs in argument 'gen'. This must contain column headers 'ps' for SNP positions/IDs, 'pvals' for

p-values and 'negLog10' for -log10(p).

logreg.bi A file name specified by either a variable of mode character, or a double-quoted

string, containing logistic regression -log10(p) for the biallelic SNPs with col-

umn names 'ps' for SNP positions/IDs and 'negLog10' for -log10(p).

logreg.tri.tetra

A file name specified by either a variable of mode character, or a double-quoted string, containing logistic regression -log10(p) for the tri and tetra allelic SNPs

with column names 'ps' for SNP positions/IDs and 'negLog10' for -log10(p).

var.matrix A vector of file names specified by double-quoted strings. The files should con-

tain presence absence matrices, with rows being variants (of 0s and 1s), and columns being samples. Column headers must contain 'ps' for variant posi-

tions/IDs with the others being the sample names.

logreg.var A vector of file names specified by double-quoted strings, of files containing lo-

gistic regression -log10(p-value) results for the presence absence matrices. Column names must contain 'ps' for variant positions/IDs and 'negLog10' for the

-log10(p).

1mm.var A vector of file names specified by double-quoted strings, of files containing

GEMMA results for the presence absence matrices.

cutOffCor Correlation cut-off for assigning and colouring variants by Principal Compo-

nents (Default = 0, variants are coloured by the PC they are most correlated

with).

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run. 1mm Whether to run GEMMA (Default = TRUE).

maf Minor allele frequency for GEMMA (Default = 0, all varaints are tested).

relmatrix A file name specified by either a variable of mode character, or a double-quoted

string of a file containing the GEMMA relatedness matrix of the samples created from biallelic SNPs. The individual ordering must be in the same order as the

column names in argument 'gen'.

lognull The log likelihood under the null from GEMMA.

lambda Lambda from GEMMA. output.dir Output file directory.

 ${\tt creating All Plots}$ 

Whether to create all bugwas plots. Default = TRUE.

allBranchAndPCCor

Whether or not to retreive correlation matrix between branches and PCs. Default = FALSE.

# **Examples**

```
lin_loc()
## An example of running lin_loc with the minimum required inputs
## Assuming gemma is installed in the present working directory
gen <- system.file("extdata", "gen.txt", package = "bugwas")
pheno <- system.file("extdata", "pheno.txt", package = "bugwas")
phylo <- system.file("extdata", "tree.txt", package = "bugwas")
prefix <- "test_bugwas"
gem.path <- "./gemma"
data <- lin_loc(gen = gen, pheno = pheno, phylo = phylo, prefix = prefix, gem.path = gem.path)</pre>
```

logregVsLMM

Plot of the P-values of logistic regression versus those of LMM

# **Description**

This function generates the plot of the P-values of logistic regression versus those of SNP GWAS.

#### Usage

```
logregVsLMM(config, biallelic, triallelic, colourPalette = NULL)
```

# **Arguments**

config A list called 'config' created from the lin\_loc function. It is a required input.

biallelic A list called 'biallelic' created from the lin\_loc function. It is a required input.

triallelic A list called 'triallelic' created from the lin\_loc function. It is a required input.

A list called 'triallelic' created from the lin\_loc function. It is a required input.

A vector of colours colour the significant principal components identified by the Bayesian Wald test (see testGenomeWidePCs). If this is NULL then colours are chosen from a default colour palette.

```
data <- lin_loc(gen = gen, pheno = pheno, phylo = phylo,
    prefix = prefix, gem.path = gem.path)
logregVsLMM(biallelic = data$biallelic, triallelic = data$triallelic, config = data$config)</pre>
```

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pcl	Load	1n	gP.	Lot

Plot PC loadings of all SNPs.

# **Description**

This function plots the loadings of all SNPs for each significant principle component identified by the Bayesian Wald test.

## Usage

```
pcLoadingPlot(config, biallelic)
```

# **Arguments**

config A list called 'config' created from the lin\_loc function. It is a required input.

biallelic A list called 'biallelic' created from the lin\_loc function. It is a required input.

# **Examples**

```
data <- lin_loc(gen = gen, pheno = pheno, phylo = phylo,
    prefix = prefix, gem.path = gem.path)
pcLoadingPlot(config = data$config, biallelic = data$biallelic)</pre>
```

plotIndividualBy2PCs Plot of the sample on the first two principal components

# Description

This function generates a plot of the sample on the first two principal components.

# Usage

```
plotIndividualBy2PCs(biallelic = NULL, config = NULL)
```

# **Arguments**

biallelic A list called 'biallelic' created from the lin\_loc function. It is a required input.

config A list called 'config' created from the lin\_loc function. It is a required input.

```
data <- lin_loc(gen = gen, pheno = pheno, phylo = phylo,
    prefix = prefix, gem.path = gem.path)
plotIndividualBy2PCs(biallelic = data$biallelic, config = data$config)</pre>
```

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snpManhattanPlot	Generates Manhattan plots for a SNP GWAS	

#### **Description**

This function generates the Manhattan plot(s) for a SNP GWAS.

# Usage

```
snpManhattanPlot(biallelic, triallelic, config, colourPalette = NULL)
```

# **Arguments**

biallelic A list called 'biallelic' created from the lin\_loc function. It is a required input. triallelic A list called 'triallelic' created from the lin\_loc function. It is a required input. A list called 'config' created from the lin\_loc function. It is a required input. config colourPalette A vector of colours colour the significant principal components identified by the Bayesian Wald test (see testGenomeWidePCs). If this is NULL then colours are

chosen from a default colour palette.

### **Examples**

```
data <- lin_loc(gen = gen, pheno = pheno, phylo = phylo,</pre>
prefix = prefix, gem.path = gem.path)
snpManhattanPlot(biallelic = data$biallelic, triallelic = data$triallelic,
 config = data$config)
```

snpPCManhattanPlot

Plot the a Manhattan plot organised by the signficance of PCs.

# **Description**

This function generates a Manhattan plot organised by the signficance of the principal components.

# Usage

```
snpPCManhattanPlot(config, biallelic, triallelic, colourPalette = NULL)
```

#### **Arguments**

config	A list called 'config' created from the lin_loc function. It is a required input.
biallelic	A list called 'biallelic' created from the lin_loc function. It is a required input.
triallelic	A list called 'triallelic' created from the lin_loc function. It is a required input.
colourPalette	A vector of colours colour the significant principal components identified by the Bayesian Wald test (see testGenomeWidePCs). If this is NULL then colours are chosen from a default colour palette.
p.genomewidepc	A matrix of the significant principal component and their correlation with lineages. This is Rayesian Wald test results produced by the function test Genome Wil

eages. This is Bayesian Wald test results produced by the function testGenomeWide-PCs. If this is NULL then testGenomeWidePCs is called to generate the required

test results.

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#### **Examples**

```
data <- lin_loc(gen = gen, pheno = pheno, phylo = phylo,
   prefix = prefix, gem.path = gem.path)
snpPCManhattanPlot(biallelic = data$biallelic, triallelic = data$triallelic,
   config = data$config)</pre>
```

testGenomeWidePCs

Testing the genome-wide principal components

# **Description**

This function performs the genome-wide principal components.

# Usage

```
testGenomeWidePCs(config = NULL, biallelic = NULL)
```

# **Arguments**

config A list called 'config' created from the bugwas function.
biallelic A list called 'biallelic' created from the bugwas function.

#### Value

The p-value of the Bayesian Wald test for the genome-wide effect of principal components.

# **Examples**

```
data <- lin_loc(gen = gen, pheno = pheno, phylo = phylo,
    prefix = prefix, gem.path = gem.path)
testGenomeWidePCs(config = data$config, biallelic = data$biallelic)</pre>
```

testGenomeWidePCsAll Testing the genome-wide principal components for all PCs.

# Description

This function identifies the non-genome-wide principal components.

# Usage

```
testGenomeWidePCsAll(config = NULL, biallelic = NULL)
```

#### **Arguments**

config A list called 'config' created from the bugwas function
biallelic A list called 'biallelic' created from the bugwas function

#### Value

The p-value of the Bayesian Wald test for the genome-wide effect of all principal components.

#### **Examples**

```
data <- lin_loc(gen = gen, pheno = pheno, phylo = phylo,
    prefix = prefix, gem.path = gem.path)
testGenomeWidePCsAll(config = data$config, biallelic = data$biallelic)</pre>
```

trueAndPredPhenoOnTreePlot

Generates all plots.

# **Description**

This function that plots the true and predicted phenotypes on a tree.

# Usage

```
trueAndPredPhenoOnTreePlot(config, biallelic, treeInfo, p.genomewidepc = NULL,
  colourPalette = NULL)
```

#### **Arguments**

config A list called 'config' created from the lin\_loc function. It is a required input.

biallelic A list called 'biallelic' created from the lin\_loc function. It is a required input.

treeInfo A list called 'treeInfo' created from the lin\_loc function. It is a required input.

p.genomewidepc A matrix of the significant principal component and their correlation with lineages. This is Bayesian Wald test results produced by the function testGenomeWidePCs. If this is NULL then testGenomeWidePCs is called to generate the required test results.

colourPalette A vector of colours colour the significant principal components identified by the Bayesian Wald test (see testGenomeWidePCs). If this is NULL then colours are chosen from a default colour palette.

```
data <- lin_loc(gen = gen, pheno = pheno, phylo = phylo,
  prefix = prefix, gem.path = gem.path)
trueAndPredPhenoOnTreePlot(biallelic = data$biallelic, treeInfo = data$treeInfo,
  config = data$config)</pre>
```

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