Package 'POPPATHR'

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Title Population-based pathway analysis of SNP-SNP coevolution

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Depends R (>= $3.5.0$)
Imports plyr, dplyr, ggplot2, data.table, stringr, reshape2, gdata, RColorBrewer, gridExtra, cowplot, GenomicRanges, snpStats, RCy3
Description A bioinformatics package for determining pathway-level SNP coevolution driven by population-based positive selection.
License GPL (>= 2)
Encoding UTF-8
URL https://github.com/rosscm/POPPATHR RoxygenNote 6.1.1
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calcFST	Calculates SNP-level FST values for use in setupGSEArun.R	

Description

Calculates SNP-level FST values for use in setupGSEArun.R

Usage

```
calcFST(genoF, famF, outDir)
```

Arguments

genoF (char) path to file with SNP genotype data (PLINK format).

famF (char) path to PLINK population coded fam file.

outDir (char) directory to store output files.

Value

none

getPathStats Generates SNP lists per selection-enriched and unenriched pathway

as determined by GSEA

Description

Generates SNP lists per selection-enriched and unenriched pathway as determined by GSEA

Usage

```
getPathStats(genoF, resF, gseaStatF, snp2geneF, realFam, enrichNES = 0.3,
  unenrichNES = 0.1, enrichDir, unenrichDir)
```

Arguments

genoF (char) path to file with SNP genotype data (PLINK format).

resF (char) path to files with GSEA results. Strutured to compare results of two

population analyses i.e., CEU vs. YRI and CEU vs. LWK.

gseaStatF (char) path to GSEA statistics file. snp2geneF (char) path to SNP-gene mapping file.

realFam (char) path to PLINK population coded fam file.

enrichNES (integer) NES cutoff to select validated selection-enriched pathways (default=0.3)

 $unenrichNES \qquad (integer) \ NES \ cutoff \ to \ select \ unenriched \ pathways \ (default=0.1)$

enrichDir (char) path to directory to store output files (PLINK files per validated selection-

enriched pathway)

unenrichDir (char) path to directory to store output files (PLINK files per unenriched path-

way)

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Value

none

LDstatsBPM Calculate selection statistics (LD) and perform exploratory analyses for two sets of variants via R snpStats package

Description

Calculate selection statistics (LD) and perform exploratory analyses for two sets of variants via R snpStats package

Usage

LDstatsBPM(enrichDir, unenrichDir, pop1, pop2, snp2geneF, outDir)

Arguments

enrichDir (char) path to selection-enriched pathway SNP lists

unenrichDir (char) path to unenriched pathway SNP lists

pop1 (char) character code for the first population (controls). pop2 (char) character code for the second population (cases).

snp2geneF (char) path to file with snp2gene mappings. Output of mapSNP2gene() (found

 $in\ GWAS2Pathway)$

outDir (char) path to output directory

Value

none

LDstatsWPM	Calculate selection statistics (LD) and perform exploratory analyses
	for two sets of variants via R snpStats package

Description

Calculate selection statistics (LD) and perform exploratory analyses for two sets of variants via R snpStats package

Usage

```
LDstatsWPM(enrichDir, unenrichDir, pop1, pop2, outDir)
```

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Arguments

enrichDir (char) path to selection-enriched pathway SNP lists

unenrichDir (char) path to unenriched pathway SNP lists

pop1 (char) character code for the first population (controls).
pop2 (char) character code for the second population (cases).

outDir (char) path to output directory

Value

none

plotEmap Create EnrichmentMap in Cytoscape to visualize selection-enriched

pathways

Description

Create a network where nodes are pathway gene sets significantly enriched in two independent population-based GSEA (i.e., validated selection-enriched gene sets.) Pathways are then clustered to identify main pathways and themes of the selection-enriched gene sets.

Usage

```
plotEmap(gmtF, eMapF, outDir, netName = "generic", imageFormat = "png",
  verbose = FALSE)
```

Arguments

gmtF (char) file path to GMT file (this needs to be absolute path name).

outDir (char) path to directory where file should be stored.

netName (char) name for network in Cytoscape. imageFormat (char) one of PNG, PDF, SVG, or JPEG.

verbose (logical) print messages.

not run because requires Cytoscape to be installed and open # plotEmap(gmtFile = gmtF, nodeAttrFile = nodeAttrFile, netName="HighRisk", # outDir=outDir).

resF (char) file path to results_forEM_selEnrich.txt (generated by writeEmapFile.R).

Value

Filename of image to which EnrichmentMap is exported. Also side effect of plotting the EnrichmentMap in an open session of Cytoscape.

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popPCA	Calculate population stratification between different ancestry groups (e.g., CEU vs YRI) via complete linkage clustering

Description

Calculate population stratification between different ancestry groups (e.g., CEU vs YRI) via complete linkage clustering

Usage

```
popPCA(genoF, famF, pop1, pop2, dimensions = 3L, outF)
```

Arguments

genoF	(char) path to file with SNP genotype data (PLINK format).
famF	(char) path to PLINK fam file (case/control population coded).
pop1	(char) character code for the first population (controls).
pop2	(char) character code for the second population (cases).
dimensions	(integer, default=3).
outF	(char) path to write PNG image.

Value

none

recodeFAM	Recodes PLINK fam file to case/control format by population adapted
	from SP plink_baseSetup.R (part of GWAS2pathway package)

Description

Recodes PLINK fam file to case/control format by population adapted from SP plink_baseSetup.R (part of GWAS2pathway package)

Usage

```
recodeFAM(genoF, pop1, pop2, popsF, setSeed = 42L, caseCode = 2,
  ctrlCode = 1, outF = "case-ctrl")
```

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Arguments

genor	(char) path to file with SNP genotype data (PLINK format)
pop1	(char) character code for the first population (controls).
pop2	(char) character code for the second population (cases).
popsF	(char) path to file with population information. Gives the number of samples per population in the dataset.
setSeed	(integer) value for set.seed() before shuffling (default=42).
caseCode	(integer) value for case samples in fam phenotype column.

(abov) mothers file middle CND construct data (DI INIV formers)

outF (char) optional - name for fam file (default=case-ctrl). file extension added.

(integer) value for control samples in fam phenotype column.

Value

none

ctrlCode

setupGSEArun	Sets up and runs GSEA using population-based FST table containing
	GSEA results (Geneset, Size, ES, NES, NominalP, FDR, FWER)

Description

Sets up and runs GSEA using population-based FST table containing GSEA results (Geneset, Size, ES, NES, NominalP, FDR, FWER)

Usage

```
setupGSEArun(realF, pathF, snp2geneF, setPerm = 1000L,
   snp2genedist = 500000L, minGene = 10L, maxGene = 300L,
   setSeed = 42L, outDir)
```

Arguments

realF (char) path to file with real SNP association statistics

pathF (char) path to pathway definitions GMT file

snp2geneF (char) path to file with snp2gene mappings. Output of mapSNP2gene() (found

 $in\ GWAS2 Pathway).$

setPerm (integer) set cycle of permutations to run default=1000

 $snp2genedist \qquad (integer) \ value \ for \ GSEA-distance. \ Max. \ distance \ between \ SNP \ and \ gene \ for$

their association in snp2geneF

minGene (integer) value for GSEA -setmin. Min. number of genes in a gene set to be

considered

maxGene (integer) value for GSEA -setmax. Max. number of genes in a gene set to be

considered

setSeed (integer) value for GSEA -seed

Value

none

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SNP2gene	Maps SNPs to their nearest genes (for use in setupGSEArun.R) (adapted from map_SNP2gene from SP's GWAS2pathway package)

Description

NOTE: updated GenomicRanges to version 1.30.1 on 01/30/2018 Requires ignore.strand=TRUE param to properly run distanceToNearest() given unknown strand assignment for SNP array-based genotyping info

Usage

```
SNP2gene(inF, geneF, marg = 0L, outDir)
```

Arguments

inF (char) path to PLINK .bim file (only CHR, SNP, and BP columns considered.

geneF (char) path to refseq table with header.

marg (integer) region upstream and downstream of [txStart,txEnd].

outDir (char) path to write snp-to-gene mapping.

Value

none

writeEmapF	Write enrichments file for use in Cytoscape EnrichmentMap

Description

Write enrichments file for use in Cytoscape EnrichmentMap

Usage

```
writeEmapF(resEmF, enrichNES = 0.3, outF)
```

Arguments

resEmF (list) file paths to results_forEM.txt for each population comparison (generated

by setupGSEA_run.R).

enrichNES (integer) NES cutoff to select validated selection-enriched pathways (default=0.3)

outF (char) name of output file.

Value

none. Output file write into directory of first population comparison. E.g., CEU_YRI directory

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