Package 'rJPSGCS'

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Description

R wrappers for gene drop programs in Alun Thomas' Java Programs for Statistical Genetics and Computational Statistics (JPSGCS).

Details

R wrappers for programs from JPSGCS required to simulate data by gene drop (GeneDrops) based on a fitted LD model (FitGMLD). JPSGCS java source code was downloaded from Alun Thomas' website (no longer active) in March 2011. See the package NEWS file (RShowDoc("NEWS", package="rJPSGCS")) for a list of modifications needed to interface with R.

Author(s)

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exdat example data for rJPSGCS

Description

Example data, comprised of simulated genotypes for 10 SNPs on the same chromosome, for five unrelated individuals, along with a genetic map for the SNPs. The fifth subject has missing genotypes at SNPs 3 and 4.

Usage

```
data(exdat)
```

Format

A list with the following elements:

\$markers a numeric matrix of SNP genotypes coded as 0, 1 or 2 copies of an index allele a numeric vector containing the genetic map positions for the SNPs in centiMorgans

Examples

```
data(exdat)
```

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FitGMLD	estimate a graphical model for linkage disequilibrium	

Description

This function is an R wrapper for the JPSGCS program to estimate a graphical model for linkage disequilibrium (LD) from a sample of genotypes. The genetic markers are assumed to be on the same chromosome.

Usage

```
FitGMLD(par = "input.par", fped = "flipped.ped", out.ld.par = "out.ld.par")
```

Arguments

par	Name of a LINKAGE parameter file (also called a DATAFILE: see http://linkage.rockefeller.edu/soft/linkage/) that contains marker and genetic map information.
fped	Name of a LINKAGE pedigree file "flipped" so that rows correspond to loci and columns correspond to individuals.
out.ld.par	Name of the file in the current working directory to contain the fitted LD model.

Details

The input pedigree file fped is not the standard format where rows correspond to individuals and columns to loci. Rather, it has been transposed so that rows correspond to loci. Transposed LINKAGE pedigree files may be written by the write.pedfile function with the option transpose=TRUE.

Further information on the graphical model is available from the article listed in the References.

Value

None. The result of the function call is the output file out.ld.par.

Note

This function can require large amounts of memory for large data sets (e.g. whole chromosomes of SNPs at the marker density of the HapMap project). The computation for fitting the LD model is done in java, using functions from Alun Thomas' suite of Java Programs for Statistical Genetics and Computational Statistics (JPSGCS). The JPSGCS java programs are accessed by R-wrappers provided by the **rJPSGCS** package, which initializes the java virtual machine (JVM) if not already done so by another package. To over-ride the default amount of memory java allocates for heap space, initialize the JVM *before* loading **rJPSGCS** as follows:

```
options(java.parameters="-Xmx2048m") #set max heap space to 2GB library(rJava)
```

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```
.jinit() #initialize the JVM
library(rJPSGCS) # now load rJPSGCS
```

Author(s)

Sigal Blay, Jinko Graham and Brad McNeney

References

Thomas A. Estimation of graphical models whose conditional independence graphs are interval graphs and its application to modeling linkage disequilibrium. Comput Stat Data Anal. 2009; 53:1818-1828.

See Also

```
write.parfile, write.pedfile
```

Examples

```
## Not run:
data(exdat)
sdat<-as(exdat$markers,"snp.matrix") #coerce to snp.matrix
# Write a LINKAGE parameter file
write.parfile(snp.data=sdat,map=exdat$map,file="test.par")
# Write a "flipped" pedfile for the SNP data on unrelated subjects.
write.pedfile(pedinf="unrelated",snp.data=sdat,file="ftest.ped",transpose=TRUE)
FitGMLD(par="test.par",fped="ftest.ped",out.ld.par="test.ld.par")
# Clean up
unlink(c("test.par","ftest.ped","test.ld.par"))
## End(Not run)</pre>
```

 ${\tt GeneDrops}$

simulate genotypes on a pedigree

Description

This function is an R wrapper for the JPSGCS multi-locus gene-drop program.

Usage

```
GeneDrops(ld.par = "ld.par", ped = "in.ped", n, complete.data=FALSE,
gd.ped = "gd.ped", compress.pedfiles=FALSE)
```

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Arguments

ld.par	Name of a LINKAGE parameter file or an LD model file output by FitGMLD.
ped	Name of the post-MAKEPED pedfile (see the explanation in the documentation for read.pedfile) giving the pedigree relationships and marker data availability to simulate.
n	Number of pedigrees to simulate.
complete.data	Should complete data be simulated; i.e., should we ignore the marker data availability in ped? Default is FALSE.
gd.ped	Name of the output file of simulated marker data in post-MAKEPED pedigree file format.
compress.pedfiles	

Should the output pedigree file be compressed? Default is FALSE.

Value

No value. The output is in the file named by gd.ped.

Note

This function can require large amounts of memory for large data sets (e.g. whole chromosomes of SNPs at the marker density of the HapMap project). The computations are done in Java, using functions from Alun Thomas' suite of Java Programs for Statistical Genetics and Computational Statistics (JPSGCS). The JPSGCS Java programs are accessed by R-wrappers provided by the **rJPSGCS** package, which initializes the Java virtual machine (JVM) if not already done so by another package. To over-ride the default amount of memory Java allocates for heap space, initialize the JVM before loading **rJPSGCS** as follows:

```
options(java.parameters="-Xmx2048m") #set max heap space to 2GB
library(rJava)
.jinit() #initialize the JVM
library(rJPSGCS) # now load rJPSGCS
```

Author(s)

Sigal Blay, Jinko Graham and Brad McNeney

See Also

FitGMLD

Examples

```
# 1. Gene-drop assuming independent loci in the founders, using allele
# frequencies and map information from the example data set.
data(exdat)
# Print out the data to examine missing data pattern.
exdat$markers
```

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```
#Coerce to snp.matrix
sdat<-as(exdat$markers,"snp.matrix")</pre>
# Need LINKAGE parameter and pedigree files as input.
# For independent loci in the founders use a standard LINKAGE parameter file.
write.parfile(snp.data=sdat, map=exdat$map, file="indep.par")
# To construct the pedigree file for unrelated individuals, use
# the pedinf="unrelated" option.
write.pedfile(pedinf="unrelated", snp.data=sdat, file="unrel.ped")
GeneDrops(ld.par="indep.par",ped="unrel.ped",n=10,gd.ped="gd.ped")
# Read the SNP data back into R.
simdat<-read.pedfile("gd.ped")$snp.data</pre>
# Look at the simulated data: need to coerce to numeric matrix to see
# values of individual genotypes, rather than the usual snp.matrix summary
as(simdat, "numeric")
# 2. Simulate data at 5 independent SNPs 0.1 cM apart on a parent-child trio.
map=(1:5)/10
# Marker data on each individual as follows:
mom=c(1,2,1,1,1) # complete data
dad=c(0,NA,0,1,0) # missing data at SNP 2
child=c(NA,1,1,2,NA) #missing data at SNPs 1 and 5
sdat<-rbind(mom,dad,child)</pre>
colnames(sdat)<-paste("SNP",1:5,sep="")</pre>
sdat<-as(sdat,"snp.matrix")</pre>
write.parfile(snp.data=sdat, map=map, file="trio.par")
# Block of code setting up pedigree information for mom, dad and child.
# See the Details section of the write.pedfile help file for
# information on the file format.
trioinf<-matrix(c(</pre>
    1, 1, 0, 0, 0, 0, 0, 1, \#mom: ID=1, no parents in pedigree
    1, 2, 0, 0, 0, 0, 1, 1, \#dad: ID=2, no parents in pedigree
    1, 3, 2, 1, 0, 0, 0, 2, 1), #child: ID=3, dadID=2, momID=1
    byrow=TRUE, ncol=9, nrow=3)
write.pedfile(pedinf=trioinf, snp.data=sdat,file="trio.ped")
GeneDrops(ld.par="trio.par",ped="trio.ped",n=10,gd.ped="gd.ped")
#Read the SNP data back into R.
simdat<-read.pedfile("gd.ped")$snp.data</pre>
# Look at the missing data patterns in simdat
as(simdat, "numeric")
## Not run:
# 3. Repeat gene drop for the trio, but this time use an LD model fit
# by FitGMLD. FitGMLD requires a LINKAGE parameter file and a transposed
# pedigree file as input. Can use the parameter file trio.par again.
# The transposed pedigree file can be generated as follows:
write.pedfile(pedinf=trioinf,snp.data=sdat,file="f.ped",transpose=TRUE)
FitGMLD("trio.par", "f.ped", "out.ld.par")
#Call to GeneDrops uses the output of FitGMLD as the parameter file and
# the regular (not transposed) pedigree file
GeneDrops(ld.par="out.ld.par",ped="trio.ped",n=10,gd.ped="gd.ped")
simdat<-read.pedfile("gd.ped")$snp.data</pre>
unlink(c("f.ped","out.ld.par"))
```

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```
## End(Not run)
unlink(c("unrel.ped","trio.ped","indep.par","trio.par", "gd.ped"))
```

read.pedfile

read a LINKAGE pedigree file into a snp.matrix object

Description

This function reads post-MAKEPED pedigree files (used by JPSGCS) into a snp.matrix object. The post-MAKEPED format includes nine columns of pedigree structure and subject characteristics (see **Details** below) before the marker data. By contrast, pre-MAKEPED pedigree file format has only six column of pedigree and subject data before the marker data. This function was adapted from the read.snps.pedfile function in the **chopsticks** package (formerly **snpMatrix**), which reads pre-MAKEPED pedigree files.

Usage

```
read.pedfile(file, snp.names=NULL, assign=NULL, missing=NULL,
X=FALSE, sep=".")
```

Arguments

•	•	
	file	The name of the pedigree file to be read.
	snp.names	A character vector of SNP names. If NULL the function will look for SNP names in a corresponding .map or .info file with the same file name root as file (e.g., if file="myfile.ped" it will look for myfile.map or myfile.info).
	assign	A list of named mappings from letters to alleles; not currently used.
	missing	Missing data code in the pedigree file; not currently used.
	Χ	Is the pedigree file comprised of X-chromosome SNPs? Default is FALSE.
	sep	Character string to separate Family and Member IDs in the row.names of the output snp.matrix object; not used.

Details

JPSGCS pedigree files are in post-MAKEPED format (though much of this information is ignored):

Column 1:	Pedigree number
Column 2:	Individual ID number
Column 3:	ID of father; 0=no father in pedigree
Column 4:	ID of mother; 0=no mother in pedigree
Column 5:	First offspring ID; ignored by JPSGCS
Column 6:	Next paternal sibling ID; ignored by JPSGCS
Column 7:	Next maternal sibling ID; ignored by JPSGCS
Column 8:	Sex; 1=male, 2=female
Column 9:	Proband status (1=proband, 0=not); ignored by JPSGCS

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The pre-MAKEPED format excludes columns 5 - 7. read.pedfile is essentially read.snps.pedfile for post-MAKEPED pedigree files, except that the option low.mem of read.snps.pedfile has not been implemented.

As in read.snps.pedfile, when reading in SNP genotype data the function looks for a corresponding .map or .info file with information on the SNPs such as physical map positions. For example, if file="test.ped", the function looks for test.map or test.info to read SNP information. Any SNP information that is found is saved in the element snp.support of the output object (see **Value**).

Value

A list with three components:

```
snp.data a snp.matrix object holding the genotypes,
subject.support
a data frame containing the first nine columns of the pedigree file, and
snp.support a data frame of SNP information. NB: this is only meaningful if a .map or .info
```

file was found.

Author(s)

Sigal Blay, Jinko Graham and Brad McNeney

See Also

```
write.pedfile, read.snps.pedfile
```

Examples

```
data(exdat)
sdat<-as(exdat$markers,"snp.matrix") #coerce to snp.matrix
#Write pedigree file for unrelated subjects
write.pedfile(pedinf="unrelated",snp.data=sdat,file="test.ped")
#Read it back into R
sdat2<-read.pedfile("test.ped")
#clean up
unlink("test.ped")</pre>
```

write.parfile

write a LINKAGE parameter file

Description

Use a snp.matrix object and genetic map to create a LINKAGE parameter file.

Usage

```
write.parfile(snp.data, map, file="out.par")
```

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Arguments

snp.data	a snp.matrix object of genotypes
map	genetic map in centiMorgans

file file name for the output parameter file

Value

None. The output is the parameter file.

Author(s)

Jinko Graham, Brad McNeney and Annick Nembot-Simo

See Also

```
write.pedfile
```

Examples

```
data(exdat)
sdat<-as(exdat$markers,"snp.matrix") #coerce to snp.matrix
write.parfile(sdat,exdat$map,file="test.par")
unlink("test.par")</pre>
```

write.pedfile

write a snp.matrix object to a LINKAGE pedigree file.

Description

Write a snp.matrix object to a LINKAGE pedigree file, with pedigree information followed by genotypes. This function can be used to write either post-MAKEPED or pre-MAKEPED pedigree file formats (see **Details**). In addition to standard pedigree file formats the function can optionally write transposed pedigree files, as required by FitGMLD.

Usage

```
write.pedfile(pedinf, snp.data, file, transpose=FALSE, sep=" ", eol="\n", na="0")
```

Arguments

pedinf The pedigree information for the pedfile as a matrix or data frame. Altherately,

users may specify pedinf="unrelated" for unrelated subjects. See the Exam-

ples.

snp.data A snp.matrix object containing the marker data.

file File name for the output pedfile.

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transpose	Should the pedigree file be transposed, as required by FitGMLD? Default is FALSE.
sep	Field separator (not currently used).
eol	The end-of-line character.
na	The missing data code in snp.data.

Details

JPSGCS pedigree files are in post-MAKEPED format (though much of this information is ignored):

```
Column 1: Pedigree number

Column 2: Individual ID number

Column 3: ID of father; 0=no father in pedigree

Column 4: ID of mother; 0=no mother in pedigree

Column 5: First offspring ID; ignored by JPSGCS

Column 6: Next paternal sibling ID; ignored by JPSGCS

Column 7: Next maternal sibling ID; ignored by JPSGCS

Column 8: Sex; 1=male, 2=female

Column 9: Proband status (1=proband, 0=not); ignored by JPSGCS
```

The pre-MAKEPED format excludes columns 5 - 7.

Value

A numeric vector comprised of the number of subjects and SNPs written to the pedigree file.

Warning

The function makes **NO** attempt to check the pedigree information for correctness. It is up to the user to specify valid pedigree information.

Author(s)

Sigal Blay, Jinko Graham and Brad McNeney

See Also

```
read.pedfile, read.snps.pedfile
```

Examples

```
data(exdat)
sdat<-as(exdat$markers,"snp.matrix") #coerce to snp.matrix
#Write a post-MAKEPED pedigree file for unrelated subjects.
pedinf<-matrix(
    c( 1, 1, 0, 0, 0, 0, 0, 1, 1,
        2, 1, 0, 0, 0, 0, 0, 1, 1,
        3, 1, 0, 0, 0, 0, 0, 1, 1,
        4, 1, 0, 0, 0, 0, 0, 1, 1,</pre>
```

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```
5, 1, 0, 0, 0, 0, 0, 1, 1),
byrow=TRUE, ncol=9, nrow=5)
write.pedfile(pedinf,sdat,file="test.ped")
#clean up
unlink("test.ped")
```

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