Package 'GenABEL'

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 $\operatorname{\mathsf{add}}.\operatorname{\mathsf{phdata}}$

Adds phenotypic variables to gwaa.data object

Description

Adds phenotypic variables to phdata part of an gwaa.data-class object

Usage

```
add.phdata(data, newphdata, name)
```

Arguments

data an object of gwaa.data-class

newphdata data frame or a vector with new phenotypic data

name if 'newphdata' is a vector, the name of new variable should be specified in

'name'

Details

If "newphdata" is a data frame, it is simply merged to the phdata part of the "data", and is sorted according to the right order. In this case, The "newphdata" frame should contain single variable named "id", preferably of "character" class. It may contain "sex" variable, but that will be re-named to avoid duplication with the default sex variable presented in phdata.

If 'newphdata' is a vector, it should be of the same length as the number of people in the 'data' and is assumed to have the same order. In this case, you also need to supply the name of the new phenotype via the 'name' argument

Value

An (updated) object of gwaa.data-class

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Author(s)

Yurii Aulchenko

See Also

```
merge.gwaa.data merge.snp.data
```

Examples

```
require(GenABEL.data)
data(srdta)
# take a small subset for this example
srdta <- srdta[1:10,1:5]</pre>
srdta
# add single var
rnd <- rnorm(nids(srdta))</pre>
srdta1 <- add.phdata(srdta,rnd,name="random")</pre>
srdta1
# add > 1 var
# generate id names
ids <- paste("p",c(2,1,7,3,5,9,11,22,27),sep="")
# generate some random trait values
newtra1 <- rnorm(9)</pre>
newtra2 <- rnorm(9)</pre>
# make data frame
x <- data.frame(id=ids,newtra1=newtra1,newtra2=newtra2)</pre>
# now add this new trait to the data
srdta1 <- add.phdata(srdta,x)</pre>
srdta1
```

add.plot

function to plot additional GWAA results

Description

Add plot of results of GWA analysis

Usage

```
add.plot(x, ..., df = 1, col=c("lightgreen", "lightblue"), sort=TRUE, delta = 1)
```

Arguments

```
x object of type scan.gwaa-class, as returned by scan.glm, qtscore, ccfast, emp.ccfast, emp.qtscore, or scan.haplo; or of type scan.gwaa.2D-class, as returned by scan.haplo.2D or scan.glm.2D.
```

.. additional arguments to be passed to plot

df	P-value at which df to add (1, 2 or "Pc1df")
col	which colors to use to depict consecutive chromosomes
sort	whether results should be plotted after sorting by chromosome and position
delta	gap width between chromosomes

Value

No value returned.

Author(s)

Yurii Aulchenko

See Also

```
plot, snp.subset, scan.glm, qtscore, ccfast, emp.qtscore, emp.ccfast, scan.haplo, scan.haplo.2D, scan.glm.2D
```

Examples

```
require(GenABEL.data)
data(srdta)
a <- ccfast("bt",srdta,snps=c(1:100))
plot(a)
a1 <- qtscore(bt,srdta,snps=c(1:100))
add.plot(a1,col="red",type="l")</pre>
```

```
arrange_probabel_phe arranges ProbABEL phenotype-file
```

Description

Function to arrange ProbABEL phenotype-file; it takes phenotypic data as input and aligns that with genotypic data of ProbABEL

Usage

```
arrange_probabel_phe(modelterms, phedata, gendata,
  file = "probabel.PHE")
```

Arguments

modelterms	vector of character, which specifies the variables to be included into ProbABEL phenotype-file. Should contain, and start with 'id' column, which should provide the same ID codes as these in gendata
phedata	phenotypic data (matrix, data.frame, or gwaa.data-class object)
gendata	genetic data to be used with ProbABEL, either databel-class object or name of the (index/data) file containing filevector data for ProbABEL
file	name of the ProbABEL phenotype file

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Value

file with phenotypes ready for use with ProbABEL

Author(s)

Yurii Aulchenko

```
as.character.gwaa.data
```

Attempts to convert genotypic part of gwaa.data to character

Description

A function to convert @gtdata slot of an object of gwaa.data-class to "character"

Usage

```
## S3 method for class 'gwaa.data'
as.character(x, ...)
```

Arguments

```
x An object of gwaa.data-class
```

Value

A matrix containing genotypes in character format

Author(s)

Yurii Aulchenko

See Also

```
as. character. snp. data, as. double. gwaa. data, as. double. snp. data, as. hsgeno, as. genotype. gwaa. data, as. genotype. snp. data
```

```
require(GenABEL.data)
data(srdta)
as.character(srdta[1:5,1:10])
```

```
as.character.snp.coding
```

Attempts to convert internal snp.coding-class to character

Description

A function to convert an object of snp.coding-class to "character"

Usage

```
## S3 method for class 'snp.coding'
as.character(x, ...)
```

Arguments

```
x An object of snp.coding-class
...
```

Value

A vector containing actual (nucleotide) coding, for corresponding SNPs, in character format

Author(s)

Yurii Aulchenko

See Also

```
as.character.snp.strand, as.character.snp.data, as.double.snp.data, as.hsgeno, as.genotype.snp.data
```

```
require(GenABEL.data)
data(srdta)
as.character(srdta@gtdata@coding[1:5])
```

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Description

A function to convert an object of snp.data-class to "character"

Usage

```
## S3 method for class 'snp.data' as.character(x, ...)
```

Arguments

```
x An object of snp.data-class
```

Value

A matrix containing genotypes in character format

Author(s)

Yurii Aulchenko

See Also

```
as.double.snp.data, as.hsgeno, as.genotype.snp.data
```

```
require(GenABEL.data)
data(srdta)
as.character(srdta@gtdata[1:5,1:10])
```

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```
as.character.snp.strand
```

Attempts to convert internal strand-class to character

Description

A function to convert an object of snp.strand-class to "character"

Usage

```
## S3 method for class 'snp.strand'
as.character(x, ...)
```

Arguments

```
x An object of snp.strand-class ...
```

Value

A vector containing strand ("+", "-" or "u"), for corresponding SNPs, in character format

Author(s)

Yurii Aulchenko

See Also

```
as.character.snp.coding, as.character.snp.data, as.double.snp.data, as.hsgeno, as.genotype.snp.data
```

```
require(GenABEL.data)
data(srdta)
as.character(srdta@gtdata@strand[1:5])
```

as.data.frame.gwaa.data

```
as.data.frame.gwaa.data

Attempts to convert snp.data to "hsgeno"
```

Description

A function taking @phdata part (data.frame) of the object of gwaa.data-class

Usage

```
## S3 method for class 'gwaa.data' as.data.frame(x, ...)
```

Arguments

```
x An object of data. frame-class
```

Details

Use is mainly internal

Value

A data-frame containing phenotypic data

Author(s)

Yurii Aulchenko

See Also

```
as.character.snp.data, as.double.snp.data, as.genotype.snp.data
```

```
require(GenABEL.data)
data(srdta)
as.data.frame(srdta[1:5,])
```

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as.double.gwaa.data Attempts to convert gwaa.data to double

Description

A function to convert an object of gwaa.data-class to "double"

Usage

```
## S3 method for class 'gwaa.data' as.double(x, ...)
```

Arguments

```
x An object of gwaa.data-class
```

Value

A matrix containing genotypes in double (numeric) format

Author(s)

Yurii Aulchenko

See Also

```
as.character.gwaa.data,as.character.snp.data,as.double.gwaa.data,as.double.snp.data,as.hsgeno,as.genotype.gwaa.data,as.genotype.snp.data
```

```
require(GenABEL.data)
data(srdta)
as.double(srdta[1:5,1:10])
```

as.double.snp.data

as.double.snp.data

Attempts to convert snp.data to double

Description

A function to convert an object of snp.data-class to "double"

Usage

```
## S3 method for class 'snp.data'
as.double(x, ...)
```

Arguments

```
x An object of snp.data-class ...
```

Value

A matrix containing genotypes in double (numeric) format

Author(s)

Yurii Aulchenko

See Also

```
as.character.snp.data, as.hsgeno, as.genotype.snp.data
```

Examples

```
require(GenABEL.data)
data(srdta)
as.double(srdta@gtdata[1:5,1:10])
```

as.genotype

Attempts to convert object to "genotype"

Description

A function to convert an object to "genotype" data frame

Usage

```
as.genotype(x, \ldots)
```

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Arguments

```
x An object of snp.data-class ...
```

Value

A data-frame containing "genotype" data class, consumable by "genetics" library

Author(s)

Yurii Aulchenko

See Also

```
as.character.gwaa.data,as.character.snp.data,as.double.gwaa.data,as.double.snp.data,as.hsgeno,as.genotype.gwaa.data,as.genotype.snp.data
```

Examples

```
require(GenABEL.data)
data(srdta)
as.genotype(srdta@gtdata[1:5,1:10])
```

```
as.genotype.gwaa.data Attempts to convert gwaa.data to "genotype"
```

Description

A function to convert @gtdata slot of an object of gwaa.data-class to "genotype" data frame

Usage

```
## S3 method for class 'gwaa.data' as.genotype(x, ...)
```

Arguments

```
x An object of gwaa.data-class ...
```

Value

A data-frame containing genotypes consumable by "genetics" library

Author(s)

Yurii Aulchenko

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See Also

```
as.character.gwaa.data,as.character.snp.data,as.double.gwaa.data,as.double.snp.data,as.hsgeno,as.genotype.gwaa.data,as.genotype.snp.data
```

Examples

```
require(GenABEL.data)
data(srdta)
as.genotype(srdta[1:5,1:10])
```

 ${\tt as.genotype.snp.data} \quad \textit{Attempts to convert snp.data to "genotype"}$

Description

A function to convert an object of snp.data-class to "genotype" data frame

Usage

```
## S3 method for class 'snp.data'
as.genotype(x, ...)
```

Arguments

```
x An object of snp.data-class
```

Value

A data-frame containing genotypes consumable by "genetics" library

Author(s)

Yurii Aulchenko

See Also

```
as.character.snp.data, as.double.snp.data, as.hsgeno
```

```
require(GenABEL.data)
data(srdta)
as.genotype(srdta@gtdata[1:5,1:10])
```

as.hsgeno.gwaa.data

as.hsgeno

Attempts to convert object to "hsgeno"

Description

A function to convert an object to "hsgeno" data frame, to be used by "haplo.stats" library

Usage

```
as.hsgeno(x, ...)
```

Arguments

```
x An object of snp.data-class or gwaa.data-class
```

Value

A data-frame containing alleles, consumable by "haplo.stats" library

Author(s)

Yurii Aulchenko

See Also

```
as.character.snp.data, as.double.snp.data, as.genotype.snp.data
```

Examples

```
require(GenABEL.data)
data(srdta)
as.hsgeno(srdta[1:5,1:3])
as.hsgeno(srdta@gtdata[1:5,1:3])
```

as.hsgeno.gwaa.data

Attempts to convert gwaa.data to "hsgeno"

Description

A function to convert @gtdata slot of an object of gwaa.data-class to "hsgeno" data frame

Usage

```
## S3 method for class 'gwaa.data'
as.hsgeno(x, ...)
```

as.hsgeno.snp.data

Arguments

```
x An object of gwaa.data-class ...
```

Value

A data-frame containing alleles, consumable by "haplo.stats" library

Author(s)

Yurii Aulchenko

See Also

```
as.character.gwaa.data,as.character.snp.data,as.double.gwaa.data,as.double.snp.data,as.hsgeno,as.genotype.gwaa.data,as.genotype.snp.data
```

Examples

```
require(GenABEL.data)
data(srdta)
as.hsgeno(srdta[1:5,1:10])
```

as.hsgeno.snp.data

Attempts to convert snp.data to "hsgeno"

Description

A function to convert an object of snp.data-class to "hsgeno" data frame

Usage

```
## S3 method for class 'snp.data'
as.hsgeno(x, ...)
```

Arguments

```
x An object of snp.data-class ...
```

Value

A data-frame containing alleles, consumable by "haplo.stats" library

Author(s)

Yurii Aulchenko

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See Also

```
as.character.snp.data, as.double.snp.data, as.genotype.snp.data
```

Examples

```
require(GenABEL.data)
data(srdta)
as.hsgeno(srdta@gtdata[1:5,1:10])
```

autosomal

Function telling all autosomal SNPs

Description

Function telling all autosomal SNPs

Usage

```
autosomal(data)
```

Arguments

data

object of gwaa.data-class or snp.data-class

Details

For every SNP, looks up the chromosome, and, when it is an utosome (not X, Y, XY, or mt), reports the name back

Value

Vector of SNP names

Author(s)

Yurii Aulchenko

```
require(GenABEL.data)
data(ge03d2)
autosomal(ge03d2)[1:10]
```

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blurGenotype	blur genotype calls into probabilites
	3 · · · · · · · · · · · · · · · · · · ·

Description

'blurs' genotype calls into probabilities: translates single genotype g2, into probability distribution P(g1|g2), that is probability that true genotype is g1 given g2 is the observed 'called' genotype and error rate is epsilon. Probability that 'true' genotype is called genotype is set to $(1-epsilon)^2$, the probability that true genotype differs at 1 allele is set to epsilon*(1-epsilon), and both allelel differ = epsilon*(1-epsilon).

Usage

```
blurGenotype(g, q = NULL, epsilon = 0.01)
```

Arguments

g	vector of genotypes for a particular person (at locus 1, locus 2, etc., coded as 0,
	1, 2 (corresponding to genotypes AA, AB, and BB, respectively) and NA.
q	(optional) vector of coded allele frequencies for locus 1, locus 2, etc.
epsilon	error rate

Value

matrix with columns corresponding to SNPs and rows corresponding to 'g0', 'g1', 'g2'. For a particular SNP, a vale in cell 'gK' is the probability that true genotype is 'K', given thw original call and error-rate.

Author(s)

Yurii Aulchenko

```
require(GenABEL.data)
data(srdta)
# select 10 first SNPs
df <- srdta[,1:10]
# compute effect allele freq
EAF <- summary(gtdata(df))$"Q.2"
EAF
# get genotypes of first 5 people
g1 <- as.numeric(df[1:5,])
g1
# blur the genotype of person 1, snp 1
blurGenotype(g1[1,1])
# blur all genotypes of person 2; assume no info for missing
blurGenotype(g1[2,])</pre>
```

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```
# blur all genotypes of person 2; use HWE to infer missing blurGenotype(g1[2,],q=EAF)
```

catable

function to generate summary table for quantitative data

Description

This function makes a table with number of observations which fall between user-defined categories

Usage

```
catable(data, categories = c(quantile(data,c(0.01,0.1,0.5,0.9,0.99),na.rm=TRUE)),
cumulative = FALSE, na.rm = TRUE, digits = 3)
```

Arguments

data A vector of numerics

categories vector containing desired cut-off levels

cumulative whether cumulative distribution should be shown

na.rm how to treat NAs

digits number of digits to be saved in rounding

Value

table with number and proportion of observations falling between categories

Author(s)

Yurii Aulchenko

See Also

```
summary.snp.data, perid.summary
```

```
require(GenABEL.data)
data(srdta)
callr <- summary(srdta@gtdata)[,"CallRate"]
catable(callr,c(0.93,0.95,0.99))
catable(callr)
catable(callr,cum=TRUE)</pre>
```

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ccfast	fast case-control analysis	

Description

Fast case-control analysis by computing chi-square test from 2x2 (allelic) or 2x3 (genotypic) tables

Usage

Arguments

У	character name of the vector of case-control status. Cases are denoted as 1 and controls as $\boldsymbol{0}$.
data	An object of gwaa.data-class
snpsubset	Index, character or logical vector with subset of SNPs to run analysis on. If missing, all SNPs from data are used for analysis.
idsubset	Index, character or logical vector with subset of IDs to run analysis on. If missing, all people from data are used for analysis.
times	If more then one, the number of replicas to be used in derivation of empirical genome-wide significance. See emp.qtscore, which calls qtscore with times>1 for details
quiet	do not print warning messages
bcast	If the argument times > 1, progress is reported once in bcast replicas
clambda	If inflation facot Lambda is estimated as lower then one, this parameter controls if the original P1df (clambda=TRUE) to be reported in Pc1df, or the original 1df statistics is to be multiplied onto this "deflation" factor (clambda=FALSE). If a numeric value is provided, it is used as a correction factor.
propPs	proportion of non-corrected P-values used to estimate the inflation factor Lambda, passed directly to the estlambda

Value

```
Object of class scan.gwaa-class
```

Author(s)

Yurii Aulchenko

See Also

```
emp.ccfast, plot.scan.gwaa, scan.gwaa-class
```

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Examples

```
require(GenABEL.data)
data(srdta)
a <- ccfast("bt",data=srdta,snps=c(1:10),ids=c(1:100))
a
a <- ccfast("bt",data=srdta)
plot(a)</pre>
```

check.marker

function to do genotypic quality control

Description

This function helps selecting the marker which should enter into GWA analysis based on call rate, minor allele frequency, value of the chi-square test for Hardy-Weinberg equilibrium, and redudndance, defined as concordance between the distributions of the genotypes (including missing values).

Usage

```
check.marker(data, snpsubset, idsubset, callrate = 0.95,
    perid.call=0.95, extr.call = 0.1, extr.perid.call = 0.1, het.fdr = 0.01,
    ibs.threshold = 0.95, ibs.mrk = 2000, ibs.exclude="both", maf, p.level = -1,
    fdrate = 0.2, odds = 1000, hweidsubset, redundant = "no",
    minconcordance = 2.0, qoption = "bh95", imphetasmissing = TRUE, XXY.call=0.8,
    intermediateXF = c(0.5, 0.5))
```

Arguments

data	gwaa.data or snp.data object
snpsubset	a subset of SNPs to check (names, indexes, logical), default is all from data
idsubset	a subset of people to check (names, indexes, logical), default is all from data
callrate	cut-off SNP call rate
perid.call	cut-off individual call rate (maximum percent of missing genotypes in a person)
extr.call	SNPs with this low call rate are dropped prior to main analysis
extr.perid.cal	1
	people with this low call rate are dropped prior to main analysis
het.fdr	FDR rate for unacceptably high individual heterozygosity
ibs.threshold	threshold value for acceptable IBS
ibs.mrk	How many random markers should be used to estimate IBS. When ibs.mrk < 1, IBS checks are turned off. When "all" all markers are used.
ibs.exclude	"both", "lower" or "none" – whether both samples with IBS>ibs.threshold should be excluded, the one with lower call rate, or no check (equivalent to use of 'ibs.mrk = -1').

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maf cut-off Minor Allele Frequency. If not specified, the default value is 5 chromo-

somes 5/(2*nids(data))

p.level cut-off p-value in check for Hardy-Weinberg Equilibrium. If negative, FDR is

pplied

fdrate cut-off FDR level in check for Hardy-Weinberg Equilibrium

odds cut-off odds to decide whether marker/person should be excluded based on

sex/X-linked marker data inconsistency

hweidsubset a subset of people to check (names, indexes, logical) to use for HWE check

redundant if "bychrom", redundancy is checked within chromosomes; "all" – all pairs of

markers; "no" - no redundancy checks

minconcordance a parameter passed to "redundant" function. If "minconcordance" is > 1.0 only

pairs of markers which are exactly the same, including NA pattern, are considered as redundant; if 0 < "minconcordance" < 1, then pairs of markers with concordance > "minconcordance" are considered redundant. see redundant for details. Note that if "minconcordance" < 1 the program will take much longer

time to run

qoption if "bh95", BH95 FDR used; if "storey", qvalue package (if installed) is used

imphetasmissing

If "impossible heterozygotes" (e.g. heterozygous mtDNA, and male Y- and X-chromsome markers) should be treated as missing genotypes in the QC proce-

dure

XXY.call What proportion of Y-chromosome markers should be called to consider that

Y-chromosome is present (in presence of XX)

intermediateXF X-chromosomal F to be considered 'intermediate' and regarded as error; cur-

rently use of default disables this check

Details

In this procedure, sex errors are identified initally and then possible residual errors are removed iteratively. At the first step, of the iterative procedure, per-marker (minor allele frequency, call rate, exact P-value for Hardy-Weinberg equilibrium) and between-marker statistics are generated and controlled for, mostly using the internal call to the function summary.snp.data.

At the second step of the iterative procedure, per-person statistics, such call rate within a person, heterozygosity and and between-person statistics (identity by state across a random sample of markers) are generated, using perid.summary and ibs functions. Heterozygosity and IBS are estimated using only autosomal data. If IBS is over ibs.threshold for a pair, one person from the pair is added to the ibsfail list and excluded from the idok list. At the second step, only the markers passing the first step are used.

The procedure is applied recursively till no further markers and people are eliminated.

Value

Object of class check.marker-class

Author(s)

Yurii Aulchenko

24 check,marker-class

See Also

check.trait,ibs, summary.snp.data,perid.summary,plot.check.marker, summary.check.marker,
redundant, HWE.show, check.marker-class

Examples

```
# usual way
require(GenABEL.data)
data(ge03d2c)
# truncate the data to make the example faster
ge03d2c \leftarrow ge03d2c[seq(from=1,to=nids(ge03d2c),by=2),seq(from=1,to=nsnps(ge03d2c),by=2)]
# many errors
mc0 <- check.marker(ge03d2c)</pre>
# take only people and markers passing QC
fixed0 <- ge03d2c[mc0$idok,mc0$snpok]</pre>
# major errors fixed, still few males are heterozygous for X-chromsome markers
mc1 <- check.marker(fixed0)</pre>
# fix minor X-chromosome problems
fixed1 <- Xfix(fixed0)</pre>
# no errors
mc2 <- check.marker(fixed1)</pre>
summary(mc2)
# ready to use fixed1 for analysis
# let us look into redundancy
require(GenABEL.data)
data(srdta)
mc <- check.marker(data=srdta,ids=c(1:300),call=.92,perid.call=.92)</pre>
names(mc)
mc$nohwe
mc <- check.marker(data=srdta@gtdata[,1:100],call=0.95,perid.call=0.9,</pre>
maf=0.02,minconcordance=0.9,fdr=0.1,redundant="all",ibs.mrk=0)
summary(mc)
HWE.show(data=srdta,snps=mc$nohwe)
plot(mc)
```

check.marker-class

Class "check.marker"

Description

This class contains results of genotypic quality control. This is an list object, usually generated by check.marker.

Names

snpok Markers which passed all criteriaidok People which passed all criteria

check.marker-class 25

```
nohwe Markers which did not pass HWE check
```

Pex.nohwe Exact HWE P-values for markers which did not pass HWE check

nocall Markers with call rate < specified callrate

nofreq Markers with MAF < specified maf

Xmrkfail X-linked markers with too many heterozygous male genotypes

redundant Redundant markers

details.redundancy List with details on redundant markers (reference-marker <-> redundant-markers)

idnocall People with too low SNP call rate across al SNPs

hetfail People having too high heterozygosity

ibsfail People having too high IBS with other people

Xidfail Men with too many heterozygous X-linked markers

call List with details on call: call, name (of marker), map, chromosome

Methods

```
summary signature(object = "check.marker"): gives a cross table summrising how many
markers did not pass because of this or that criteria

plot signature(object = "check.marker"): Plots summary of genotypic data QC
```

Author(s)

Yurii Aulchenko

See Also

```
check.marker, summary.check.marker, redundant, plot.check.marker
```

```
require(GenABEL.data)
data(srdta)
mc <- check.marker(data=srdta@gtdata[,1:100],redundant="all",maf=0.01,
minconcordance=0.9,fdr=.1,ibs.mrk=0)
class(mc)
names(mc)
names(mc)
names(mc$call)
mc$nohwe
mc$Pex.nohwe
summary(mc)
plot(mc)</pre>
```

26 check.trait

check.trait

function to do primitive trait quality control

Description

This function check for outliers (using FDR framework) and plots the raw data.

Usage

```
check.trait(trait, data, fdrate = 0.05, graph = TRUE, binshow = FALSE,
qoption = "bh95")
```

Arguments

trait	name (or list of names) of trait(s) to be checked
data	gwaa.data object or data frame containing the trait
fdrate	false discovery rate to apply for QC
graph	if graphical output should be produced
binshow	if binary traits should be plotted
qoption	how to compute q-values (not implemented, currently using only BH95)

Details

The P-value that a particular measurement is an outlier is compted as following. Consider trait vector Y with particular i^{th} measurement denode as y_i . Let Y(-i) is vector, which is the same as Y, except that i^{th} measurement is dropped. Then Chi-square for measurement is computed as

$$Chi_i = (mean(Y(-i)) - y_i)^2 / var(Y(-i))$$

P-value is computed using 1 d.f., and the vector of P-values enters FDR computation procedure (BH95 by default).

Value

No value returned, output is made to the screen and graphical device.

Author(s)

Yurii Aulchenko

See Also

check.marker

Examples

```
require(GenABEL.data)
data(srdta)
check.trait("qt3",data=srdta)
n <- names(srdta@phdata)
check.trait(n,data=srdta)</pre>
```

checkPackageVersionOnCRAN

checks what is the version of package on CRAN

Description

Checks what is the version of package on CRAN. The CRAN page (baseUrlCRAN+packageName) is checked and parsed extracting the line with "Package source: packageName_Version.tar.gz" e.g. "Package source: GenABEL_1.6-9.tar.gz" and then the 'Version' is returned. Otherwise, NULL is returned.

Usage

```
checkPackageVersionOnCRAN(packageName,
  baseUrlCRAN = "http://cran.r-project.org/web/packages/",
  timeout = 2)
```

Arguments

packageName name of the package to check baseUrlCRAN path to CRAN repository timeout web chack timeout

Value

string containing CRAN version of the package

Author(s)

Yurii Aulchenko

```
library(GenABEL)
packageVersion("GenABEL")
checkPackageVersionOnCRAN("GenABEL")
```

28 cocohet

cocohet

Test for compound heterozygote effects

Description

Detecting rare recessive and compound heterozygote alleles in genome wide association.

Usage

```
cocohet(data, trait, window, return_all_result = TRUE,
  makePlot = FALSE, test = "CHI2",
  min_expected_cut_off = -1)
```

Arguments

data Genotype data for analysis. Object of class snp. data

trait Vector with binary trait data. Object of class integer or numeric.

window Number of SNPs on the "right" of a given SNP which are used in analysis with

a SNP. Object of class integer

return_all_result

If FALSE then return only a vector where each element is a chisq obtained as a maximum chisq between a given SNP and SNPs on the right within a window. If TRUE then return also a matrix where chisq's for all tests are stored. Object

of class logical

makePlot whether the Manhattan-type plot should be produced (TRUE or FALSE)

test Name of the test to be performed. Available tests are "CHI2", "YATES" (chi2

with Yates correction), and "FISHER". Object of class character

min_expected_cut_off

In case this is >=0 and test is NOT Pearson's chisq test then Pearson's chisq test (!) is performed only for SNPs which produce acontingency table where the expected number of subjects in each field is >min_expected_cut_off. Otherwise the specified test is performed. Object of class integer or numeric

Details

The function is an inplementation of the method aimed to detect a gene-phenotype association caused by recessive and compound heterozygote genotype states of multiple rare variants at a particular gene locus. This method is described in 'Detecting Low Frequent Loss-of-Function Alleles in Genome Wide Association Studies with Red Hair Color as Example'; Fan Liu, Maksim V. Struchalin, Kate van Duijn, Albert Hofman, Andre G. Uitterlinden, Yurii S. Aulchenko, and Manfred Kayser. PLoS ONE 6(11): e28145. doi:10.1371/journal.pone.0028145

The three tests are implemented: Pearson's chi-square test, Pearson's chi-square test with Yates correction, Fisher exact test. In case when the input parameter min_expected_cut_off is <0 the choosen in the input parameter "test" test is performed. If min_expected_cut_off >= 0 then always Pearson's chi-square test is performed exept of the cases when expected number of subjects in a field of contingency table is <min_expected_cut_off. In this case the test choosen in the input parameter test is performed.

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Value

A list is returned.

Author(s)

Maksim Struchalin

References

Fan Liu, Maksim V. Struchalin, Kate van Duijn, Albert Hofman, Andre G. Uitterlinden, Yurii S. Aulchenko, and Manfred Kayser. Detecting Low Frequent Loss-of-Function Alleles in Genome Wide Association Studies with Red Hair Color as Example'. PLoS ONE 6(11): e28145. doi:10.1371/journal.pone.0028145

Examples

```
require(GenABEL.data)
data(srdta)
chis2_nocorrection <- cocohet(data=gtdata(srdta),
trait=phdata(srdta)$bt, window=3, test="CHI2")</pre>
```

convert.snp.affymetrix

function to convert genotypic data from Affymetrix to internal format

Description

Converts genotypic data from Affymetrix format to internal genotypic data formated file

Usage

```
convert.snp.affymetrix(dir, map, outfile, skipaffym)
```

Arguments

dir Directory which affymetrix files storages.

map File name with map (annotation) information.

outfile Output data file.

skipaffym Number of lines to skip in the Affymetrix file.

Details

```
Affymetrix file has following format:
```

some information...

some information...

some information...

SNPID Call Confidence others column ...

AFFX-7317060 AB 0.01709367 ...

AFFX-7317061 BB 0.01683776 ...

AFFX-7317067 AB 0.01704767 ...

AFFX-7317077 AB 0.01817814 ...

AFFX-7317078 AA 0.0006741961 ...

AFFX-7317079 AA 0.004776776 ...

AFFX-7317063 AB 0.006349149 ...

AFFX-7317064 AB 0.04771883 ...

AFFX-7317067 AA 0.04387166 ...

The first several lines do not contain genotype information and have to be skiped. Skiped numbers of lines can be setted. by setting skipaffym input parameter. For above examle it has to be skipaffym=3.

Every row corresponds to a SNP. The first column is snp name, the second - genotype. The second column can contain letters (AA, AB, BB) or figures (1, 2, 3). Another values consider as unmeasured.

All affymetrix files must have same SNP amount and same SNP order.

The first two lines in the map file will be skiped.

If SNP does not exist in map (annotation) file this SNP will be skiped.

Output will be writen into file pointed in outfile.

Value

Does not return any value, but writes file with GenABEL raw data

Note

The function does not check if "outfile" already exists, thus it is always over-written

Author(s)

Maksim Struchalin

See Also

load.gwaa.data,convert.snp.text,convert.snp.mach,convert.snp.tpedconvert.snp.illumina

convert.snp.illumina 31

Examples

```
## Not run:
convert.snp.affymetrix(dir="where_is_our_aff_files", map="map_file",
outfile="output.raw", skipaffym=3)
## End(Not run)
```

 ${\tt convert.snp.illumina} \begin{tabular}{ll} \textit{function to convert genotypic data from Illumina/Affymetrix to internal} \\ \textit{format} \end{tabular}$

Description

Converts genotypic data from Illumina/Affymetrix-like format to internal genotypic data formated file

Usage

```
convert.snp.illumina(infile, outfile, strand = "+", bcast = 10000000)
```

Arguments

infile	Map + genotypic data file name
outfile	Output data file
strand	Specification of strand, one of "u" (unknown), "+", "-" or "file". In the latter
	case, extra column specifying the strand (again, one of "u", "+", or "-") should

bcast Reports progress after reading bcast portion of SNP genotypes

be included on the infile.

Details

Input file is the one which could be typically obtained from Illumina BeadStudio software. For example:

```
Name Chr Pos id1 id2 id3
rs1001 2 12897 AC AA AA
rs2401 3 12357 AG GG AG
rs123 3 5327 TC TT CC
```

Here, every row corresponds to a SNP, and each column, starting with the 4th, corresponds to a person.

When strand information is available (option strand="file"), the file should look like

Accepted allele codes: 1/2, A/B, A/T, A/G, A/C, T/G, T/C, G/C, A/-, T/-, G/-, C/-. Here, "-" stands of a deletion. Missing data can be coded as "-" or "00". Make sure that the coding for missing is "00" if you use one of the codings A/-, T/-, G/-, C/-!

Name Chr Pos Strand id1 id2 id3

32 convert.snp.mach

```
rs1001 2 12897 + AC AA AA
rs2401 3 12357 - AG GG AG
rs123 3 5327 + TC TT CC
Accepted strand coding: +, -, u (unknown)
```

The procedure always codes genotypes that "0", "1" and "2" correspond to AA, AB, and BB, where B is the less frequent allele. Thus GWA analysis procedures will return effect of the minor allele.

Value

Does not return any value, but writes file with GenABEL raw data

Note

The function does not check if "outfile" already exists, thus it is always over-written

Author(s)

Yurii Aulchenko

See Also

```
load.gwaa.data, convert.snp.text, convert.snp.mach, convert.snp.tped
```

Examples

```
#
# convert.snp.illumina(infile="pedin.18",out="genos.raw",strand="+")
#
```

convert.snp.mach

function to convert genotypic data from MACH format to internal data format

Description

Converts genotypic data from MACH format to internal genotypic data formated file (NOT recommended)

Usage

```
convert.snp.mach(pedfile, mapfile, infofile, outfile, quality = 0.9,
column.quality = 7, strand = "+", ...)
```

convert.snp.mach 33

Arguments

pedfile	File with genotypic data from MACH (geno or mlgeno)
mapfile	Name of the map file (note that header line should be included)
infofile	Name MACH info-file
outfile	Output data file
quality	Drop the SNPs with quality (as specified in some column of info-file) lower than this threshold.
column.quality	What column of the info-file provides "quality". Default = 7 os r2; possible values include 6 (average postrior probability).
strand	Specification of strand, one of "u" (unknown), "+", "-" or "file". In the latter case, map-file should contain an extended map (the one including strand and coding). See options to convert.snp.ped for details.
	Other arguments passed to convert.snp.ped

Details

This is a simple script converting the MACH data with convert.snp.ped, re-loading data, and filtering the snp.data object based on quality as specified in MACH info-file

Note that it is NOT recommended to process imputations results with this procedure, as uncertainty inherent to imputations is lost after processing ("hard" calls are made)! For the purposes of GWA analysis, we recommend procedures, which make direct use of estimated allele dose or genotypic probabilities.

Value

Does not return any value, but writes file with GenABEL raw data

Note

The function does not check if "outfile" already exists, thus it is always over-written

Author(s)

Yurii Aulchenko

See Also

```
load.gwaa.data,convert.snp.illumina,convert.snp.text,convert.snp.ped,convert.snp.tped
```

```
#
# convert.snp.mach(ped="pedin.18",map="map.18",out="genos.raw")
#
```

34 convert.snp.ped

convert.snp.ped	function to convert genotypic data in pedigree fromat (+map) to inter- nal data format

Description

Converts genotypic data in a variety of pedigree fromats (+map) to internal genotypic data formated file

Usage

```
convert.snp.ped(pedfile, mapfile, outfile, format = "premakeped", traits = 1,
strand = "u", bcast = 10000000, wslash=FALSE, mapHasHeaderLine=TRUE)
```

Arguments

pedfile	Pre-makeped linkage genotypic data file name (no header line)
mapfile	Name of the map file (note that by default header line should be included, see 'mapHasHeaderLine' to modify that)
outfile	Output data file
format	Input data format, either "premakeped" (default, also works with Merlin files), or "mach"
traits	How many traits are specified in the pedigree file (usually 1 – affection – or 2 – affection and liability). Has no effect when format = "mach".
strand	Specification of strand, one of "u" (unknown), "+", "-" or "file". In the latter case, map-file should contain an extended map (the one including strand and coding)
bcast	Reports progress after reading bcast portion of SNPs
wslash	Whether alleles are separated with slash (is true for Mach/Merlin format), otherwise it is assumed that alleles are separated with space. When wslash=T it is assumed that genotypes are coded with single characters, separated with slash (no spaces), e.g. "A/G", and not "A/G" or "A/G".

mapHasHeaderLine

Whether map-file has a header line

Details

Pedfile must be standard pre-makeped/Merlin linkage file, or a Mach file. In pre-makeped linkage file, columns are

ped id fa mo sex trait snp1.allele1 snp1.allele2 snp2.allele1 snp2.allele2 ...

For example

```
1 1 0 0 1 2 A A G T ...
1 2 0 0 1 0 A G T T ...
```

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```
1 3 0 0 2 1 A A T T ...
```

...

Would imply that persons 1, 2 and 3 are "founders" (which would be typical for a case-control study), 1 and 2 are males and 3 is female. Person 1 is homozygous for allele 1 at locus 1 and heterozygous at locus 2. Person 2 is heterozygous at both loci. Person 3 is homozygous for allele 2 at locus 1 and allele 1 at locus 2.

Only the second and the marker columns are used, thus make sure the IDs are unique!

Accepted allele codes: 1/2, A/B, A/T, A/G, A/C, T/G, T/C, G/C, A/-, T/-, G/-, C/-. Here, "-" stands of a deletion.

The map file is standard Merlin map. For example:

chrom name position

18 rs679153 2859916

18 rs9965482 2860891

Says that locus 1 is named rs679153 and located at chromosome 18 position 2859916. Locus 2 (rs9965482) is located at chromosome 18, position 2860891.

In extended map format, there should be 4th column specifying the strand

chrom name position strand

18 rs679153 2859916 -

18 rs9965482 2860891 +

Accepted strand coding: +, -, u (unknown)

Please note that by default the header line (e.g. "chrom name position") SHOULD be present in your file, though you can use 'mapHasHeaderLine' argument to modify this behavior

Value

Does not return any value, but writes file with GenABEL raw data

Note

The function does not check if "outfile" already exists, thus it is always over-written

Author(s)

Yurii Aulchenko

See Also

```
load.gwaa.data,convert.snp.illumina,convert.snp.mach,convert.snp.text,convert.snp.tped
```

```
#
# convert.snp.ped(ped="pedin.18",mapfile="map.18",out="genos.raw")
#
```

36 convert.snp.text

convert.snp.text	function to convert integer genotypic data file to raw internal data
	formated file

Description

Converts integer genotypic data file to raw internal data formated file

Usage

```
convert.snp.text(infile, outfile, bcast = 10000)
```

Arguments

infile Input data file name outfile Output data file

bcast Reports progress after reading bcast portion of SNPs

Details

Input genotypic data file contains all kind of genetic information. The first line of this file contains IDs of all study subjects. The second line gives names of all SNPs in the study. The third line list the chromosomes the SNPs belong to. Sequential numbers are used for autosomes and "X" (capital!) is used for the sex-chromosome. The forth line lists genomic position of the SNPs, in order which is the same as order in the line 2. The genomic position can be chromosome-specific (each chromosome starts with "0") or, better, a true genomic position (chromosome 1 starts with 0 and chromosome 2 continues at the point chromosome 1 ends).

Starting with the line five, genetic data are presented. The 5th line contains the data for SNP, which is listed first on the second line. The first column of this line specifies the genotype for the person, who is listed first on the line 1; the second column gives the genotype for the second person, so on. The genotypes are coded as 0 (missing), 1 (for AA), 2 (for AB) and 3 (for BB). Here is a small example:

```
289982 325286 357273 872422 1005389

SNP-1886933 SNP-2264565 SNP-2305014

1 1 1

825852 2137143 2585920

3 3 3 3 2

3 2 3 3 3

2 2 1 1 1
```

In this example, we can see that SNP-2305014 (number 3 in the second line) is located on chromosome 1 at the position 2585920. If we would like to know what is genotype of person with ID 325286 (second in the first line), we need to take second column and the third line of the genotypic data. This cell contains 1, thus, person 325286 has genotype "AA" at SNP-2305014.

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In the event that you do not want to use a map for some reason (such as prior ordering of the polymorphisms in the genotype file), make a dummy map-line, which contains order information.

The above described genotypic data file is (more or less) human-readable; actually, to achieve the aim of effective data storage GWAA package uses internal format. In this format, four genotypes are stored in single byte; "raw" data format of R is used.

Value

Does not return any value

Note

The function does not check if "outfile" already exists, thus it is always over-written

Author(s)

Yurii Aulchenko

See Also

```
load.gwaa.data, convert.snp.illumina, convert.snp.ped, convert.snp.mach, convert.snp.tped
```

Examples

```
#
# convert.snp.text("genos.dat","genos.raw")
#
```

convert.snp.tped

function to convert genotypic data in transposed-ped format (.tped and .tfam) to internal genotypic data formatted file

Description

Converts genotypic data in transposed-ped format (.tped and .tfam) to internal genotypic data formatted file

Usage

```
convert.snp.tped(tpedfile, tfamfile, outfile, strand = "u", bcast = 10000)
```

38 convert.snp.tped

Arguments

tpedfile Name of transposed-ped format (.tped) file to read

tfamfile Name of individual data (.tfam) file to read

outfile Name for output data file

strand Specification of strand, one of "u" (unknown), "+", "-" or "file". In the latter

case, extra column specifying the strand (again, one of "u", "+", or "-") should

be included on the tpedfile.

bcast Reports progress every time this number of SNPs have been read

Details

The transposed-ped file format may be preferred when extremely large numbers of markers have been genotyped. This file format is supported by plink! See http://pngu.mgh.harvard.edu/~purcell/plink/ for details.

The conversion is performed by C++ code that is both fast and memory efficient.

The genotype data are stored in the main transposed-ped format file, usually with a .tped file extension. If there are NSNP markers genotyped in NIND individuals, this file has NSNP rows and 4+NIND*2 columns. There is one row per marker, and no header. The first four columns are:

Chromosome

Marker name (e.g. rs number)

Genetic position (in Morgans)

Physical position (in bp)

These are followed by two columns per individual, which contain the genotype, coded as two characters. The '0' character is used for missing data. For example, a file containing data for six individuals genotyped at two SNPs would look like:

1 rs1234 0 5000650 A A 0 0 C C A C C C C C

1 rs5678 0 5000830 G T G T G G T T G T T T

In this example, the second individual is missing data for SNP rs1234, etc. The alleles can be coded by any two distinct characters, e.g. 'C' and 'G', or '1' and '2'. The '0' character is reserved for missing data, and each individual genotype must be either complete, or completely missing. In the current implementation, only the physical positions of the SNPs are read, and the genetic positions are ignored.

The indices for the columns are stored in a separate file, usually with a .tfam file extension. Traditionally, this file has six columns, and no header. In the current implementation, only the second column is used. This column must contain the individual id. Other columns are ignored.

Value

Does not return any value

Note

The function does not check if "outfile" already exists, thus it is always over-written

crnames 39

Author(s)

Toby Johnson toby.johnson@unil.ch

See Also

```
convert.snp.ped, convert.snp.illumina, convert.snp.text, convert.snp.mach, load.gwaa.data
```

Examples

```
#
# convert.snp.tped("c21.tped",map="c21.tfam",out="c21.raw")
#
```

crnames

Return column and row names

Description

Given a dimnames, returns column and row names for index cells

Usage

```
crnames(dnames,idx)
```

Arguments

dnames object dimnames

idx index (or logical condition on the original object)

Examples

```
require(GenABEL.data)
data(ge03d2ex)
a <- as.numeric(ge03d2ex[1:20,1:3])
crnames(dimnames(a),a==1)</pre>
```

40 descriptives.marker

del.phdata	delete phenotypes from phdata

Description

This function is used to delete certain phenotypes from phenotypic part (phdata) of an object of gwaa.data-class

Usage

```
del.phdata(data, what, all = FALSE)
```

Arguments

data an object of gwaa.data-class

what which phenotypes (variables) to delete, expressed as (vector of) names (charac-

ter) or integer (column of phdata data frame)

all if 'all'=TRUE and 'what' is misisng, all phenotypes are deleted, and only the

'id' and 'sex' are kept

Author(s)

Yurii Aulchenko

Examples

```
require(GenABEL.data)
data(srdta)
phdata(srdta)[1:5,]
srdta <- del.phdata(srdta,"qt1")
phdata(srdta)[1:5,]
srdta <- del.phdata(srdta,all=TRUE)
phdata(srdta)[1:5,]</pre>
```

descriptives.marker

Function to generate descriptive summary tables for genotypic data

Description

Function to generate descriptive summary tables for genotypic data

Usage

```
descriptives.marker(data,snpsubset,idsubset,file,mafc,hwec,snpc,idcc,digits = 3)
```

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Arguments

data	an object of snp.data-class or gwaa.data-class
snpsubset	Index, character or logical vector with subset of SNPs to run analysis on. If missing, all SNPs from data are used for analysis.
idsubset	Index, character or logical vector with subset of IDs to run analysis on. If missing, all people from data are used for analysis.
file	A string specifying the name of a file to write the tables to (default is missing).
mafc	vector containing desired cut-off levels for minor allele frequency
hwec	vector containing desired cut-off levels for exact HWE P-values
snpc	vector containing desired cut-off levels for SNP call rate
idcc	vector containing desired cut-off levels for individual SNP call rate
digits	number of digits to be printed

Value

A list containing descriptive tables and statistics

Author(s)

Yurii Aulchenko

Examples

```
require(GenABEL.data)
data(srdta)
descriptives.marker(srdta)
```

descriptives.scan

Function to describe "top" hits in GWA scan

Description

Describes "top" hits in GWA scan

Usage

```
descriptives.scan(data,file,top=10,sortby="P1df", sep ="\t")
```

Arguments

data	an object of snp.data-class or gwaa.data-class
file	A string specifying the name of a file to write the tables to (default is no file output).
top	How many "top" hits to describe
sortby	How to pick up "top" hits ("P1df","P2df","Pgw1df","Pgw2df")
sep	field separator (takes effect only if file argument provided)

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Value

A descritive table

Author(s)

Yurii Aulchenko

Examples

```
require(GenABEL.data)
data(srdta)
x <- qtscore(qt2,srdta)
descriptives.scan(x)</pre>
```

descriptives.trait

Function to generate descriptive summary tables for phenotypic data

Description

Function to generate descriptive summary tables for phenotypic data

Usage

```
descriptives.trait(data, subset, file, by.var=NULL, digits = 3)
```

Arguments

data	an object of snp.data-class or gwaa.data-class
subset	Subset of people to run analysis on. If missing, all people from data are used for analysis.
file	A string specifying the name of a file to write the tables to (default is no file otput).
by.var	a binary variable or a character scalar specifying the name of a binary trait in data; statistics will be produced separately for the groups and compared
digits	number of digits to be printed

Value

A table with descriptive statistics. Ptt: t-test; Pkw: kruskal.test; Pex: Fisher exact test (for factors with <5 levels)

Author(s)

Yurii Aulchenko

dprfast 43

Examples

```
require(GenABEL.data)
data(srdta)
descriptives.trait(srdta)
descriptives.trait(srdta,by.var=srdta@phdata$sex)
descriptives.trait(srdta,by.var="sex")
attach(phdata(srdta))
descriptives.trait(srdta,by.var=sex)
detach(phdata(srdta))
```

dprfast

Estimates D' between multiple markers

Description

Given a set of SNPs, computes a matrix of D'

Usage

```
dprfast(data, snpsubset, idsubset)
```

Arguments

data object of snp.data-class

snpsubset Index, character or logical vector with subset of SNPs to run analysis on. If

missing, all SNPs from data are used for analysis.

idsubset Index, character or logical vector with subset of IDs to run analysis on. If miss-

ing, all people from data are used for analysis.

Details

The function is based on slightly modified code of Hao et al.

Value

A (Nsnps X Nsnps) matrix giving D' values between a pairs of SNPs above the diagonal and number of SNP genotype measured for both SNPs below the diagonal

Author(s)

Yurii Aulchenko

References

Hao K, Di X, Cawley S. (2006) LdCompare: rapid computation of single- and multiple-marker D' and genetic coverage. Bioinformatics, 23: 252-254.

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See Also

rhofast

Examples

```
require(GenABEL.data)
data(ge03d2)
# D's using D'fast
a <- dprfast(ge03d2,snps=c(1:10))
## Not run:
library(genetics)
# D's using package genetics
b <- LD(as.genotype(ge03d2[,1:10]))$"D'"
# see that the D's are not exactly the same
cor(a[upper.tri(a)],b[upper.tri(b)])
plot(a[upper.tri(a)],b[upper.tri(b)])
## End(Not run)</pre>
```

egscore

Fast score test for association, corrected with PC

Description

Fast score test for association between a trait and genetic polymorphism, adjusted for possible stratification by principal components.

Usage

```
egscore(formula, data, snpsubset, idsubset,
  kinship.matrix, naxes = 3, strata, times = 1,
  quiet = FALSE, bcast = 10, clambda = TRUE, propPs = 1)
```

Arguments

formula	Formula describing fixed effects to be used in analysis, e.g. $y \sim a + b$ means that outcome (y) depends on two covariates, a and b. If no covariates used in analysis, skip the right-hand side of the equation.
data	An object of gwaa.data-class
snpsubset	Index, character or logical vector with subset of SNPs to run analysis on. If missing, all SNPs from data are used for analysis.
idsubset	Index, character or logical vector with subset of IDs to run analysis on. If missing, all people from data/cc are used for analysis.
kinship.matrix	kinship matrix, as returned by ibs, Use weight="freq" with ibs and do not forget to repalce the diagonal with Var returned by hom, as shown in example!
naxes	Number of axes of variation to be used in adjustment (should be much smaller than number of subjects)

egscore 45

strata	Stratification variable. If provieded, scores are computed within strata and then added up.
times	If more then one, the number of replicas to be used in derivation of empirical genome-wide significance.
quiet	do not print warning messages
bcast	If the argument times > 1, progress is reported once in bcast replicas
clambda	If inflation facot Lambda is estimated as lower then one, this parameter controls if the original P1df (clambda=TRUE) to be reported in Pc1df, or the original 1df statistics is to be multiplied onto this "deflation" factor (clambda=FALSE). If a numeric value is provided, it is used as a correction factor.
propPs	proportion of non-corrected P-values used to estimate the inflation factor Lambda, passed directly to the estlambda

Details

The idea of this test is to use genomic kinship matrix to first, derive axes of genetic variation (principal components), and, second, adjust both trait and genotypes onto these axes. Note that the diagonal of the kinship matrix should be replaced (default it is 0.5*(1+F), and for EIGENSTRAT one needs variance). These variances are porduced by hom function (see example).

The traits is first analysed using LM and with covariates as specified with formula and also with axes of variation as predictors. Corrected genotypes are defined as residuals from regression of genotypes onto axes (which are orthogonal). Correlaton between corrected genotypes and phenotype is computed, and test statistics is defined as square of this correlation times (N - K - 1), where N is number of genotyped subjects and K is the number of axes.

This test is defined only for 1 d.f.

Value

Object of class scan.gwaa-class

Author(s)

Yurii Aulchenko

References

Price A. L. et al, Principal components analysis corrects for stratification in genome-wide association studies. Nat Genet 38: 904-909.

See Also

qtscore, mmscore, ibs, scan.gwaa-class

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Examples

```
require(GenABEL.data)
data(ge03d2c)
#egscore with stratification
gkin <- ibs(ge03d2c[,autosomal(ge03d2c)],w="freq")
#replace the diagonal with right elements
diag(gkin) <- hom(ge03d2c[,autosomal(ge03d2c)])$Var
a <- egscore(dm2~sex+age,data=ge03d2c,kin=gkin)
plot(a,df="Pc1df")</pre>
```

egscore.old

Fast score test for association, corrected with PC

Description

Fast score test for association between a trait and genetic polymorphism, adjusted for possible stratification by principal components.

Usage

```
egscore.old(formula,data,snpsubset,idsubset,kinship.matrix,naxes=3,strata,
times=1,quiet=FALSE,bcast=10,clambda=TRUE,propPs=1.0)
```

Arguments

formula	Formula describing fixed effects to be used in analysis, e.g. $y \sim a + b$ means that outcome (y) depends on two covariates, a and b. If no covariates used in analysis, skip the right-hand side of the equation.
data	An object of gwaa.data-class
snpsubset	Index, character or logical vector with subset of SNPs to run analysis on. If missing, all SNPs from data are used for analysis.
idsubset	Index, character or logical vector with subset of IDs to run analysis on. If missing, all people from data/cc are used for analysis.
kinship.matrix	kinship matrix, as returned by ibs, (use weight="freq"!)
naxes	Number of axes of variation to be used in adjustment (should be much smaller than number of subjects)
strata	Stratification variable. If provieded, scores are computed within strata and then added up.
times	If more then one, the number of replicas to be used in derivation of empirical genome-wide significance.
quiet	do not print warning messages
bcast	If the argument times > 1, progress is reported once in bcast replicas

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clambda If inflation facot Lambda is estimated as lower then one, this parameter controls

if the original P1df (clambda=TRUE) to be reported in Pc1df, or the original 1df statistics is to be multiplied onto this "deflation" factor (clambda=FALSE). If a

numeric value is provided, it is used as a correction factor.

propPs proportion of non-corrected P-values used to estimate the inflation factor Lambda,

passed directly to the estlambda

Details

The idea of this test is to use genomic kinship matrix to first, derive axes of genetic variation (principal components), and, second, adjust both trait and genotypes onto these axes.

The traits is first analysed using LM and with covariates as specified with formula and also with axes of variation as predictors. Corrected genotypes are defined as residuals from regression of genotypes onto axes (which are orthogonal). Correlaton between corrected genotypes and phenotype is computed, and test statistics is defined as square of this correlation times (N - K - 1), where N is number of genotyped subjects and K is the number of axes.

This test is defined only for 1 d.f.

Value

Object of class scan.gwaa-class

Author(s)

Yurii Aulchenko

References

Price A. L. et al, Principal components analysis corrects for stratification in genome-wide association studies. Nat Genet 38: 904-909.

See Also

```
qtscore, mmscore, ibs, scan.gwaa-class
```

Examples

```
#data(ge03d2ex)
##egscore.old with stratification
#gkin <- ibs(ge03d2ex,w="freq")
#a <- egscore.old(dm2~sex+age,data=ge03d2ex,kin=gkin)
#plot(a,df="Pc1df")</pre>
```

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emp.ccfast	Genome-wide significance for a case-control GWA scan	

Description

Genome-wide significance for a case-control GWA scan. Analysis function is ccfast.

Usage

```
emp.ccfast(y, data, snpsubset, idsubset, times = 200, quiet=FALSE,
bcast = 10)
```

Arguments

data

snpsubset

idsubset

times

quiet bcast All arguments are the same as in and passed intact to the ccfast. See help for this function.

character name of the vector of case-control status. Cases are denoted as 1 and controls as 0.

An object of gwaa.data-class

Index, character or logical vector with subset of SNPs to run analysis on. If missing, all SNPs from data are used for analysis.

Index, character or logical vector with subset of IDs to run analysis on. If missing, all people from data are used for analysis.

If more then one, the number of replicas to be used in derivation of empirical genome-wide significance. See emp.qtscore, which calls qtscore with times>1 for details

do not print warning messages

If the argument times > 1, progress is reported once in bcast replicas

Details

In the analysis of empirical significance, first time the function ccfast is called and result object is saved. Later, the function ccfast is called times times with replace=FALSE in order to generate the distribution under the null. Each call, minimal P-value is extracted and compared with original P-values. For a particular SNP, empirical P-value is obtained as a proportion of times minimal Ps from resampled data was less then the original P.

The list elements effB, effAB and effBB are the ones obtained from the analysis of the original (not permuted) data set

Value

Object of class scan.gwaa-class

Author(s)

Yurii Aulchenko

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See Also

```
ccfast, emp.qtscore, scan.gwaa-class
```

Examples

```
require(GenABEL.data)
data(srdta)
a<-ccfast("bt",data=srdta,snps=c(500:800))
plot(a)
# this does not make sense, as the whole experiment must be analysed, not a small region!
# also, times = 20 is way too small (should be at least 200)
b<-emp.ccfast("bt",data=srdta,snps=c(500:800),bcast=10, times = 20)
plot(b)
# compare qvalues and empirical P
qv<-qvaluebh95(a[,"P1df"])$qval
qv
b[,"P1df"]
plot(qv,b[,"P1df"],xlim=c(0,1),ylim=c(0,1))
abline(a=0,b=1)</pre>
```

emp.qtscore

Genome-wide significance for a GWA scan

Description

Genome-wide significance for a GWA scan. Analysis function is qtscore.

Usage

```
emp.qtscore(formula , data, snpsubset, idsubset, strata, trait.type="gaussian",
times = 200, quiet=FALSE, bcast = 10)
```

Arguments

All arguments are the same as in and passed intact to the qtscore. See help for this function.

Formula describing fixed effects to be used in analysis, e.g. $y \sim a + b$ means that outcome (y) depends on two covariates, a and b. If no covariates used in analysis, skip the right-hand side of the equation.

detaula An object of gwaa.data-class

snpsubset Index, character or logical vector with subset of SNPs to run analysis on. If

missing, all SNPs from data are used for analysis.

idsubset Index, character or logical vector with subset of IDs to run analysis on. If miss-

ing, all people from data/cc are used for analysis.

strata Stratification variable. If provieded, scores are computed within strata and then

added up.

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trait.type	"gaussian" or "binomial". If not specified, the procedure quesses the type
times	If more then one, the number of replicas to be used in derivation of empirical genome-wide significance. See $emp.qtscore$, which calls $qtscore$ with times>1 for details

quiet do not print warning messages

bcast If the argument times > 1, progress is reported once in bcast replicas

Details

In the analysis of empirical significance, first time the function qtscore is called and result object is saved. Later, the function qtscore is called times times with replace=FALSE in order to generate distribution under the null. Each call, minimal P-value is extracted and compared with original P-values. For a particular SNP, empirical P-value is obtained as a proportion of times minimal Ps from resampled data was less then original P.

The list elements effB, effAB and effBB are the ones obtained from the analysis of the original (not permuted) data set

The function does not yet implement correct analysis for X-linked data.

Value

```
Object of class scan.gwaa-class
```

Author(s)

Yurii Aulchenko

See Also

```
qtscore, emp.ccfast, scan.gwaa-class
```

Examples

```
require(GenABEL.data)
data(srdta)
a<-qtscore(qt3~age+sex,data=srdta,snps=c(1:200))
plot(a)
# this does not make sense, as the whole experiment must be analysed, not a small region!
# also, times = 20 is way too small (should be at least 200)
b<-emp.qtscore(qt3~age+sex,data=srdta,snps=c(1:200), times = 20)
plot(b)</pre>
```

estlambda 51

estlambda Estimate the inflation factor for a distribution of P-values	estlambda	Estimate the inflation factor for a distribution of P-values	
--	-----------	--	--

Description

Estimate the inflation factor for a distribution of P-values or 1df chi-square test. The major use of this procedure is the Genomic Control, but can also be used to visualise the distribution of P-values coming from other tests. Methods implemented include 'median' (median(chi2)/0.455...), regression (of observed onto expected) and 'KS' (optimizing the chi2.1df distribution fit by use of Kolmogorov-Smirnov test)

Usage

```
estlambda(data, plot = FALSE, proportion = 1,
  method = "regression", filter = TRUE, df = 1, ...)
```

Arguments

data	A vector of reals. If all are <=1, it is assumed that this is a vector of P-values, else it is treated as a vector of chi-squares
plot	Whether the plot should be shown or not (default).
proportion	The proportion of lowest P (or χ^2) values to be used when estimating the inflation factor λ .
method	"regression" (default), "median", or "KS": method to be used for λ estimation.
filter	if the test statistics with 0-value of χ^2 should be excluded prior to estimation of λ .
df	Number of degrees of freedom.
	arguments passed to the plot function.

Value

A list with elements

estimate Estimate of λ

se Standard error of the estimate

Author(s)

Yurii Aulchenko

See Also

```
ccfast, qtscore
```

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Examples

```
require(GenABEL.data)
data(srdta)
pex <- summary(gtdata(srdta))[,"Pexact"]
estlambda(pex, plot=TRUE)
estlambda(pex, method="regression", proportion = 0.95)
estlambda(pex, method="median")
estlambda(pex, method="KS")
a <- qtscore(bt,srdta)
lambda(a)</pre>
```

export.impute

function to export GenABEL data in IMPUTE format

Description

Exports GenABEL data to IMPUTE/SNPTEST/GTOOLS format

Usage

```
export.impute(data,genofile="impute.gen",samplefile="impute.sample",
strandfile="impute.strand",cachesizeMb=128)
```

Arguments

data gwaa.data object

genofile Output genotype data file name

samplefile Output sample information file name, 'id' header line + IDs of people will be

written there

strandfile name for strand output file

cachesizeMb approximate amount of RAM to be used to generate chunks of data

Details

The most interesting part is the genotype file which is generated, this file contains one SNP per row, with columns

```
SNP_name_1 SNP_name_2 Position Allele1 Allele2 P_1_11 P_1_12 P_1_22 P_2_11 P_2_12 P_2_22
```

where SNP_name_1 = SNP_name_2 is SNP name, Position is SNP position, Allele1 and Allele2 are reference and effective alleles; P_i_jk is the probability that person i has genotype jk; these are (1 0 0) for genotype Allele1Allele1, (0 1 0) for genotype Allele1Allele2, (0 0 1) for genotype Allele2Allele2, and (0 0 0) if genotype is missing

Strand file contains the SNP name, position and strand, as generated by as.character(data@gtdata@strand); this is up to user if this slot contains correct data (see eg convert.snp.illumina on how to import strand info)

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Value

No value returned; all infor stored in files

Author(s)

Yurii Aulchenko

See Also

```
export.merlin.
```

Examples

```
## Not run:
require(GenABEL.data)
data(srdta)
export.impute(srdta[1:50,1:3])
## End(Not run)
```

export.merlin

function to export GenABEL data in merlin format

Description

Exports GenABEL data to Merlin and other pedigree formats

Usage

```
export.merlin(data, pedfile = "merlin.ped", datafile = "merlin.dat",
    mapfile = "merlin.map", format = "merlin", fixstrand = "no",
    extendedmap = TRUE, traits = 1, order = TRUE, stepids = 100,
    dpieceFun = "new")
```

Arguments

data	gwaa.data object	
pedfile	Output pedigree data file name	
datafile	Output data (information) file name; no output if NULL	
mapfile	Output map file name	
format	Output format, either "merlin" or "plink"	
fixstrar	"no" – the strand information and coding comes from the data; "+" – change a coding to "+" strand, "-" – change all coding to "-" strand	ıll
extended	if TRUE extended map (+ strand, + coding) is saved with the name "mapfile.ext where "mapfile" is the parameter supplied by user	.",
traits	How many fake traits to insert before first column of marker data	

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order Should output be ordered by chromosome and position

stepids make this larger for faster processing and smaller for lower RAM use

dpieceFun function used to dump data. Do use default.

Details

The use is straightforward, with only the "fixstrand" option requiring some explanation. Consider a SNP on "-" strand with alleles G and A. If this SNP is accessed on "+" strand, the corresponding alleles would be C and T. While for example Affymetrix reports SNPs on bot "+" and "-" strands, HapMap reports coding on "+" strand only. To make data compatible, and/or to run imputations, one will need to convert all SNP codes to "+" strand. This can be achieved by running export.merlin() with fixstrand="+" parameter.

Value

No value returned; files saved on HDD

Author(s)

Yurii Aulchenko

See Also

To load the data to GenABEL again, use convert.snp.ped, load.gwaa.data.

Examples

```
#
# load(srdta)
# export.merlin(srdta[1:50,1:3])
#
```

export.plink

Export GenABEL data in PLINK format

Description

Export GenABEL data in PLINK format. This function is a simple wrapper to the export.merlin function with specific arguments + few lines of code to export phenotypes

Usage

```
export.plink(data, filebasename = "plink",
  phenotypes = "all", transpose = TRUE,
  export012na = FALSE, ...)
```

Arguments

data GenABEL data object of 'gwaa.data'-class to be exported.

filebasename base file name for exported data, extensions '.ped', '.map' and '.phe' (for phe-

notype file) are added for specific output files.

phenotypes NULL (no phenotypes exported), "all" (default) for all phenotypes or a vector

of character with names of phenotypes to be exported.

transpose if TRUE (default), 'tped' files will be produced, else 'ped' files are produced.

export012na if TRUE, export in numeric (0, 1, 2, NA) format, as opposed to ATGC format

(default: FALSE).

... arguments passed to export.merlin.

Author(s)

Yurii Aulchenko

```
extract.annotation.impute
```

extracts SNP annotation from IMPUTE files

Description

This function extracts SNP annotation information from IMPUTE files. The major problem at the moment that info-file format of IMPUTE is a little bit unstable (reported information and column order varies between impute v1, v2, and beta-version). Therefore take special care to read specification of 'order_info_snp_pos_freq1_info_qual_type'

Usage

```
extract.annotation.impute(genofile, infofile,
  chromosome = NA, order_geno_snp_a0_a1 = c(2, 4:5),
  skip_geno = 0,
  order_info_snp_pos_freq1_info_qual_type = c(2:7),
  skip_info = 1, allow_duplicated_names = FALSE)
```

Arguments

genofile IMPUTE genotype file name infofile IMPUTE info-file name

chromosome chromosome order_geno_snp_a0_a1

which columns to extract from geno-file, and what is the order for snp name, a0,

and a0? (default is OK)

skip_geno how many lines of geno-file are to be skept? (default is OK)

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order_info_snp_pos_freq1_info_qual_type

which columns to extract from info-file, and what is the order for SNP name, position, frequency of allele 1, info (Rsq), and quality (average max post prob)? Dafult works for IMPUTE v2.0, but has to be changed for other versions. Al-

ways check!

skip_info how many lines of info-file are to be skept before information starts? IMPUTE

v2.0 has a header line, therefore skip_info=1 works fine; this may be different

for other versions of IMPUTE

allow_duplicated_names

if duplicated SNP names are allowed (same order in geno and info- files is as-

sumed then)

Value

data frame containing annotaton

Author(s)

Yurii Aulchenko

extract.annotation.mach

extracts SNP annotation from MACH/HapMap legend files

Description

This function extracts SNP annotation from MACH info and HapMap legend files.

Usage

```
extract.annotation.mach(infofile, legendfile,
  chromosome = NA)
```

Arguments

infofile MACH (ml)info-file name legendfile HapMap legend file name

chromosome chromosome

Value

data frame containing annotaton

Author(s)

Yurii Aulchenko

findRelatives 57

iduals		
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Description

This function guesses relationships (expressed as estimated number meiotic connection(s)) using genomic data. Compared to guessing relations from genomic kinship matrix, this procedure offers several enhancements:

Usage

```
findRelatives(gtdata, nmeivec = c(1:2), q = NULL,
    epsilon = 0.01, quiet = FALSE, OddsVsNull = 1000,
    OddsVsNextBest = 100, twoWayPenalty = log(10),
    doTwoWay = TRUE, vsIDs = NULL, gkinCutOff = NULL,
    kinshipMatrix = NULL)
```

Arguments

gtdata	genotypic data, either 'gwaa.data' or 'snp.data' class, or matrix or 'databel' matrix (see details for format).
nmeivec	vector providing the degree of relationship to be tested (1: parent-offspring; 2: sibs, grandparent-grandchild; etc.).
q	vector of effect allele frequencis for the data.
epsilon	genotyping error rate
quiet	if TRUE, screen outputs supressed
OddsVsNull	threshold used in relationships inferences (see details)
OddsVsNextBest	threshold used in relationships inferences (see details)
twoWayPenalty	penalty on likelihoods resulting from models assuming two meiotic pathways
doTwoWay	or not
vsIDs	specific IDs to be tested vs others
gkinCutOff	if not null, sets a threshold used to pre-screen pairs before guessing relations. If value < 0 provided, procedure sets threshold automatically (recommended)
kinshipMatrix	(genomic) kinship matrix (used if gkinCutOff!=NULL)

Details

- (1) by use of IBD/IBS 3-state space, it allows to distinguish between some pairs, which have the same kinship (e.g. parent-offspring from brother-sister; uncle-nephew from grandparent-grandchild, etc.)
- (2) it reports likelihood, allowing for more rigorous inferences

If 'gtdata' are provided as a matrix (or 'databel' matrix), genotypes should be coded as 0, 1, or 2; each SNP corresponds to a column and each ID is a row. 'q' corresponds to the frequency of 'effect' (aka 'coded') allele, which is also equivalent to the mean(SNP)/2.0 provided coding is correct.

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'nmeivec' is a sequence of integers, e.g. c(1,2) will test for parent-offspring pairs and pairs separated by two meioses (sibs, grandparent-grandchild, etc.). If 'nmeivec' does not contain '0' as its first element, it will be automatically added (testing for twins). Also, nmeivec will be updated with c(nmeivec,max(nmeivec)+1,100) to allow for thesting of testing vs. 'null' (unrelated, 100) and 'most distant' specified by user (max(nmeivec)).

While one may be interested to test only a sub-set of the data for relationships, it is recommended to provide 'q' estimated using all data available (see example).

'gkinCutOff' allows use of genomic kinship matrix (computed internally) to pre-screen pairs to be tested. Use of this option with value '-1' is recommended: in this case threshod is set to 0.5^(max(nmeivec)+2). If not NULL, only pairs passing gkinCutOff are tested with the likelihood procedure.

After likelihood estimation, inference on relationship is made. Releationship (in terms of number of meioses) is 'guessed' if odds of likelihoods under the meiotic distance providing max likelihood and under the 'null' (maximal meiotic distance tested + 100) is greater than 'OddsVsNull' parameter AND odds max-lik vs. the next-best meiotic distance is greater than 'OddsVsNextBest' parameter.

Value

A list with elements call – details of the call; profile – table detailing likelihood for all pairs tested; estimatedNmeioses – nids x nids matrix containing maximum likelihood estimate of meiotic distance for all pairs of individuals guess – same as estimatedNmeioses, but all estimates not passing inference criteria (OddsVsNull, OddsVsNextBest) are NAed; compressedGuess – same as above, but removing cows and cols with missing-only elemnts

Examples

```
require(GenABEL.data)
data(ge03d2.clean)
df <- ge03d2.clean[,autosomal(ge03d2.clean)]</pre>
df <- df[,sort(sample(1:nsnps(df),1000))]</pre>
eaf <- summary(gtdata(df))$"Q.2"
### donotrun
## Not run:
relInfo <- findRelatives(df[27:30,],q=eaf)</pre>
relInfo
# look only for 1st and 2nd degree relatives
relInfo1 <- findRelatives(df[27:30],q=eaf,gkinCutOff=-1,nmeivec=c(1,2,3))</pre>
relInfo1
relInfoVS <- findRelatives(df[27:30,],q=eaf,nmeivec=c(1:6),vsIDs=idnames(df[27:30,])[1:2])
relInfoVS
## End(Not run)
### end norun
```

formetascore 59

Description

Function to run GWA analysis – using all functions available in GenABEL – and produce output oriented for future meta-analysis

Usage

```
formetascore(formula, data, stat = qtscore, transform = "no",
build = "unknown", verbosity = 1, ...)
```

Arguments

formula	standard formula
data	object of gwaa.data-class
stat	which GWA analysis function to apply. Could be ${\tt mlreg},$ qtscore, ${\tt mmscore},$ grammar, egscore, etc.
transform	Which trait transform to apply, could be ${\tt ztransform}$ or ${\tt rntransform}$. Default value is "no" – no transformation.
build	if you need that in output, specify genomic build here (e.g. "35")
verbosity	how much output is produced? Possible values are 0, 1, and 2
	further arguments, passed to the "stat" GWA analysis function

Details

This function should be applied to analysis of quantitative traits, if meta-analusis is aimed afterwards.

A transformation is applied to the formla-defined residual, and the resulting trait is analysed with specified function. Results are arranged as data-frame.

Value

Data frame, containing GWA summary. The fields include: (1) SNP name (2) chromosome (3) position (4) number of people with available data (5) effect of the allele (6) standard error of the effect (7) P-value for the test (8) corrected P-value (we will use Genomic Control) (9) coding, with reference allele coming first (10) strand (11) frequency of the reference allele (12) Exact P-value for HWE test, etc. (depends on "verbosity" parameter.

Author(s)

Yurii Aulchenko

See Also

ztransform,qtscore

GASurv GASurv

Examples

```
require(GenABEL.data)
data(ge03d2c)
x <- formetascore(bmi ~ sex+age,ge03d2c)
x[1:10,]
x <- formetascore(bmi ~ sex+age,ge03d2c,trans=ztransform)
x[1:10,]
x <- formetascore(bmi ~ sex+age,ge03d2c,trans=rntransform,verbosity=2)
x[1:10,]</pre>
```

GASurv

Makes survival data object for reg.gwaa

Description

Helper to mlreg: makes survival data object

Usage

```
GASurv(fuptime, status)
```

Arguments

fuptime Follow-up time

status (1=event, 0=censored)

Value

Matrix with column 1 = follow-up time, and 2 = status

Author(s)

Yurii Aulchenko

See Also

mlreg

GenABEL GWAS in R

Description

GenABEL: an R package for Genome Wide Association Analysis

Details

Genome-wide association (GWA) analysis is a tool of choice for identification of genes for complex traits. Effective storage, handling and analysis of GWA data represent a challenge to modern computational genetics. GWA studies generate large amount of data: hundreds of thousands of single nucleotide polymorphisms (SNPs) are genotyped in hundreds or thousands of patients and controls. Data on each SNP undergoes several types of analysis: characterization of frequency distribution, testing of Hardy-Weinberg equilibrium, analysis of association between single SNPs and haplotypes and different traits, and so on. Because SNP genotypes in dense marker sets are correlated, significance testing in GWA analysis is preferably performed using computationally intensive permutation test procedures, further increasing the computational burden.

To make GWA analysis possible on standard desktop computers we developed GenABEL library which addresses the following objectives:

- (1) Minimization of the amount of rapid access memory (RAM) used and the time required for data transactions. For this, we developed an effective data storage and manipulation model.
- (2) Maximization of the throughput of GWA analysis. For this, we designed optimal fast procedures for specific genetic tests.

Embedding GenABEL into R environment allows for easy data characterization, exploration and presentation of the results and gives access to a wide range of standard and special statistical analysis functions available in base R and specific R packages, such as "haplo.stats", "genetics", etc.

To see (more or less complete) functionality of GenABEL, try running demo(ge03d2).

Other demo of interest could be run with demo(srdta). Depending on your user priveleges in Windows, it may well not run. In this case, try demo(srdtawin).

The most important functions and classes are:

For converting data from other formats, see

convert.snp.illumina (Illumina/Affymetrix-like format). This is our preferred converting function, very extensively tested. Other conversion functions include: convert.snp.text (conversion from human-readable GenABEL format), convert.snp.ped (Linkage, Merlin, Mach, and similar files), convert.snp.mach (Mach-format), convert.snp.tped (from PLINK TPED format), convert.snp.affymetrix (BRML-style files).

For converting of GenABEL's data to other formats, see export.merlin (MERLIN and MACH formats), export.impute (IMPUTE, SNPTEST and CHIAMO formats), export.plink (PLINK format, also exports phenotypic data).

To load the data, see load.gwaa.data.

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For conversion to DatABEL format (used by ProbABEL and some other GenABEL suite packages), see impute2databel, impute2mach, mach2databel.

For data managment and manipulations see merge.gwaa.data, merge.snp.data, gwaa.data-class, snp.data-class, snp.names, snp.subset.

For merging extra data to the phenotypic part of gwaa.data-class object, see add.phdata.

For traits manipulations see ztransform (transformation to standard Normal), rntransform (rank-transformation to normality), npsubtreated (non-parametric routine to "impute" trait's values in these medicated).

For quality control, see check.trait, check.marker, HWE.show, summary.snp.data, perid.summary, ibs, hom.

For fast analysis function, see scan.gwaa-class, ccfast, qtscore, mmscore, egscore, ibs, r2fast (estimate linkage disequilibrium using R2), dprfast (estimate linkage disequilibrium using D'), rhofast (estimate linkage disequilibrium using 'rho')

For specific tools facilitating analysis of the data with stratification (population stratification or (possibly unknown) pedigree structure), see qtscore (implements basic Genomic Control), ibs (computations of IBS / genomic IBD), egscore (stratification adjustment following Price et al.), polygenic (heritability analysis), polygenic_hglm (another function for heritability analysis), mmscore (score test of Chen and Abecasis), grammar (grammar, grammar-gc, and garmmar-gamma tests of Aulchenko et al., Amin et al., and Svishcheva et al.).

For functions facilitating construction of tables for your manuscript, see descriptives.marker, descriptives.trait, descriptives.scan.

For functions recunstructing relationships from genomic data, see findRelatives, reconstructNPs.

For meta-analysis and related, see help on formetascore.

For link to WEB databases, see show.ncbi.

For interfaces to other packages and standard R functions, also for 2D scans, see scan. glm, scan. glm. 2D, scan. haplo, scan. haplo. 2D, scan. gwaa-class, scan. gwaa. 2D-class.

For graphical facilities, see plot. scan. gwaa, plot. check. marker.

Author(s)

Yurii Aulchenko et al. (see help pages for specific functions)

References

If you use GenABEL package in your analysis, please cite the following work:

Aulchenko Y.S., Ripke S., Isaacs A., van Duijn C.M. GenABEL: an R package for genome-wide association analysis. Bioinformatics. 2007 23(10):1294-6.

If you used polygenic, please cite

Thompson EA, Shaw RG (1990) Pedigree analysis for quantitative traits: variance components without matrix inversion. Biometrics 46, 399-413.

If you used environmental residuals from polygenic for qtscore, or used grammar, please cite for original GRAMMAR

GenABEL 63

Aulchenko YS, de Koning DJ, Haley C. Genomewide rapid association using mixed model and regression: a fast and simple method for genome-wide pedigree-based quantitative trait loci association analysis. Genetics. 2007 177(1):577-85.

for GRAMMAR-GC

Amin N, van Duijn CM, Aulchenko YS. A genomic background based method for association analysis in related individuals. PLoS ONE. 2007 Dec 5;2(12):e1274.

for GRAMMAR-Gamma

Svischeva G, Axenovich TI, Belonogova NM, van Duijn CM, Aulchenko YS. Rapid variance components-based method for whole-genome association analysis. Nature Genetics. 2012 44:1166-1170. doi:10.1038/ng.2410

for GRAMMAR+ transformation

Belonogova NM, Svishcheva GR, van Duijn CM, Aulchenko YS, Axenovich TI (2013) Region-Based Association Analysis of Human Quantitative Traits in Related Individuals. PLoS ONE 8(6): e65395. doi:10.1371/journal.pone.0065395

If you used mmscore, please cite

Chen WM, Abecasis GR. Family-based association tests for genome-wide association scans. Am J Hum Genet. 2007 Nov;81(5):913-26.

For exact HWE (used in summary.snp.data), please cite:

Wigginton G.E., Cutler D.J., Abecasis G.R. A note on exact tests of Hardy-Weinberg equilibrium. Am J Hum Genet. 2005 76: 887-893.

For haplo.stats (scan.haplo, scan.haplo.2D), please cite:

Schaid DJ, Rowland CM, Tines DE, Jacobson RM, Poland GA. Score tests for association between traits and haplotypes when linkage phase is ambiguous. Am J Hum Genet. 2002 70:425-434.

For fast LD computations (function dprfast, r2fast), please cite:

Hao K, Di X, Cawley S. LdCompare: rapid computation of single- and multiple-marker r2 and genetic coverage. Bioinformatics. 2006 23:252-254.

If you used npsubtreated, please cite

Levy D, DeStefano AL, Larson MG, O'Donnell CJ, Lifton RP, Gavras H, Cupples LA, Myers RH. Evidence for a gene influencing blood pressure on chromosome 17. Genome scan linkage results for longitudinal blood pressure phenotypes in subjects from the framingham heart study. Hypertension. 2000 Oct;36(4):477-83.

See Also

DatABEL, genetics, haplo.stats, qvalue

Examples

```
## Not run:
demo(ge03d2)
demo(srdta)
demo(srdtawin)
## End(Not run)
```

generateOffspring

simulates offspring's genotypes

Description

Simulat offspring's genotypes given genotypes of the parents. No LD is assumed.

Usage

```
generateOffspring(g1, g2, q = NULL)
```

Arguments

g1	vector of genotypes of parent 1, coded with 0, 1, 2 and NA
g2	vector of genotypes of parent 2, coded with 0, 1, 2 and NA
q	vector of the frequencies of effects alleles (used to simulate missings in parental
	genotypes). If NULL, $c(0.25,0.5,0.25)$ is used.

Value

a vector containing simulated genotypes of offspring

Author(s)

Yurii Aulchenko

Examples

```
eaf <- runif(10)
g1 <- rbinom(10,2,eaf)
g2 <- rbinom(10,2,eaf)
generateOffspring(g1,g2)</pre>
```

getLogLikelihoodGivenRelation

computes logLik of two blurGenotypes

Description

Compute logLik of genotypes of person 1 given genotypes of person 2 and assumed relation between the two persons (expressed with transition probability matrix; as returned with 'makeTransitionMatrix').

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Usage

```
getLogLikelihoodGivenRelation(bGenotype1, bGenotype2,
  TransitionMatrix, q)
```

Arguments

bGenotype1 blurred genotype of person 1
bGenotype2 blurred genotype of person 2
TransitionMatrix
transition probability matrix
q vector of effect allele frequencies

Author(s)

Yurii Aulchenko

Examples

```
require(GenABEL.data)
data(srdta)
# select 10 first SNPs
df <- srdta[,1:10]
# compute effect allele freq
EAF <- summary(gtdata(df))$"Q.2"
# get genotypes of first 2 people
g1 <- as.numeric(df[1:2,])
# blur all genotypes of person 1; use HWE to infer missing
bg1 <- blurGenotype(g1[1,],q=EAF)
# blur all genotypes of person 2; use HWE to infer missing
bg2 <- blurGenotype(g1[2,],q=EAF)
# generate sib-sib transision matrices
trss <- makeTransitionMatrix(EAF,nmei=c(2,2))
getLogLikelihoodGivenRelation(bg1,bg2,trss,EAF)</pre>
```

grammar

GRAMMAR test for association in samples with genetic structure

Description

Fast approximate test for association between a trait and genetic polymorphisms, in samples with genetic sub-structure (e.g. relatives). The function implements several varieties of GRAMMAR ('gamma', 'gc', and 'raw').

Usage

```
grammar(polyObject, data,
  method = c("gamma", "gc", "raw"), propPs = 1, ...)
```

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Arguments

polyObject object returned by polygenic function

data object of gwaa.data-class

method to be used, one of 'gamma', 'gc', or 'raw'

propPs proportion of non-corrected P-values used to estimate the inflation factor Lambda,

passed directly to the estlambda

... arguments passed to the function used for computations, (qtscore)

Details

With 'raw' argument, the original GRAMMAR (Aulchenko et al., 2007) is implemented. This method is conservative and generates biased estimates of regression coefficients.

With 'gc' argument, the GRAMMAR-GC (Amin et al., 2007) is implemented. This method solves the conservativity of the test, but the Genomic Control (GC) lambda is by definition "1" and can not serve as an indicator of goodness of the model; also, the estimates of regression coefficients are biased (the same as in 'raw' GRAMMAR).

GRAMMAR-Gamma (default 'gamma' argument) solves these problems, producing a correct distribution of the test statistic, interpretable value of GC Lambda, and unbiased estimates of the regression coefficients. All together, the default 'gamma' method is recommended for use.

Value

Object of scan.gwaa-class

Author(s)

Gulnara Svischeva, Yurii Aulchenko

References

GRAMMAR-Raw: Aulchenko YS, de Koning DJ, Haley C. Genomewide rapid association using mixed model and regression: a fast and simple method for genomewide pedigree-based quantitative trait loci association analysis. Genetics. 2007 Sep;177(1):577-85.

GRAMMAR-GC: Amin N, van Duijn CM, Aulchenko YS. A genomic background based method for association analysis in related individuals. PLoS One. 2007 Dec 5;2(12):e1274.

GRAMMAR-Gamma: Svischeva G, Axenovich TI, Belonogova NM, van Duijn CM, Aulchenko YS. Rapid variance components-based method for whole-genome association analysis. Nature Genetics. 2012 44:1166-1170. doi:10.1038/ng.2410

See Also

polygenic, mmscore, qtscore

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Examples

```
# Using clean ge03d2 data
require(GenABEL.data)
data(ge03d2.clean)
# take only a small piece for speed
ge03d2.clean <- ge03d2.clean[1:200,]</pre>
# estimate genomic kinship
gkin <- ibs(ge03d2.clean[,sample(autosomal(ge03d2.clean),1000)], w="freq")</pre>
# perform polygenic analysis
h2ht <- polygenic(height ~ sex + age, kin=gkin, ge03d2.clean)
h2ht$est
# compute mmscore stats
mm <- mmscore(h2ht, data=ge03d2.clean)
# compute grammar-gc
grGc <- grammar(h2ht, data=ge03d2.clean, method="gc")</pre>
# compute grammar-gamma
grGamma <- grammar(h2ht, data=ge03d2.clean, method="gamma")</pre>
# compare lambdas
lambda(mm)
estlambda(mm[,"chi2.1df"])
lambda(grGamma)
estlambda(grGamma[,"chi2.1df"])
lambda(grGc)
estlambda(grGc[,"chi2.1df"])
# compare top results
summary(mm)
summary(grGamma)
summary(grGc)
```

gwaa.data-class

Class "gwaa.data"

Description

This class contains objects holding all GWAA data – phenotypes, genotypes and other relevant information

Slots

```
phdata: dataframe with phenotypic data used in GWAA gtdata: object of class snp.data-class used to store genotypic data, map, etc.
```

Methods

```
[ signature(x = "gwaa.data", i = "ANY", j = "ANY", drop = "ANY"): subset operations. x[i,j] will select people listed in i and SNPs listed in j.
```

show signature(object = "gwaa.data"): shows both parts of the object. Take care that the objects are usually very large!

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```
summary signature(object = "gwaa.data"): Calls standard summary to describe phenotypic
    part and calls summary.snp.data to snp.data-class

gtdata signature(object = "gwaa.data"): returns genotypic data (object of snp.data-class)

phdata signature(object = "gwaa.data"): returns phenotypic data (a data.frame)

phdata<- signature(object = "gwaa.data"): method do modify phenotypic data (a data.frame)</pre>
```

Author(s)

Yurii Aulchenko

See Also

```
snp.data-class, load.gwaa.data, snp.mx-class
```

Examples

```
require(GenABEL.data)
data(srdta)
srdta@phdata[1:10,]
srdta@gtdata[1:10,1:12]
srdta[1:10,1:12]
as.numeric(srdta@gtdata[1:12,1:10])
# very long output:
summary(srdta)
```

hom

function to compute average homozygosity within a person

Description

This function computes average homozygosity (inbreeding) for a set of people, across multiple markers. Can be used for Quality Control (e.g. contamination checks)

Usage

```
hom(data, snpsubset, idsubset, snpfreq, n.snpfreq = 1000)
```

Arguments

data	Object of gwaa.data-class or snp.data-class
snpsubset	Subset of SNPs to be used
idsubset	People for whom average homozygosity is to be computed
snpfreq	when option weight="freq" used, you can provide fixed allele frequencies
n.snpfreq	when option weight="freq" used, you can provide a vector supplying the number of people used to estimate allele frequencies at the particular marker, or a fixed number

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Details

Homozygosity is measured as proportion of homozygous genotypes observed in a person. Inbreeding for person i is estimated with

$$f_i = \frac{(O_i - E_i)}{(L_i - E_i)}$$

where O_i is observed homozygosity, L_i is the number of SNPs measured in individual i and

$$E_i = \sum_{j=1}^{L_i} (1 - 2p_j(1 - p_j) \frac{T_{Aj}}{T_{Aj} - 1})$$

where T_{Aj} is the number of measured genotypes at locus j; T_{Aj} is either estimated from data or provided by "n.snpfreq" parameter (vector). Allelic frequencies are either estimated from data or provided by the "snpfreq" vector.

This measure is the same as used by PLINK (see reference).

The variance (Var) is estimated as

$$V_i = \frac{1}{N} \sum_k \frac{(x_{i,k} - p_k)^2}{(p_k * (1 - p_k))}$$

where k changes from 1 to N = number of SNPs, $x_{i,k}$ is a genotype of ith person at the kth SNP, coded as 0, 1/2, 1 and p_k is the frequency of the "+" allele.

Only polymorphic loci with number of measured genotypes >1 are used with this option.

This variance is used as diagonal of the genomic kinship matrix when using EIGENSTRAT method. You should use as many people and markers as possible when estimating inbreeding/variance from marker data.

Value

A matrix with rows corresponding to the ID names and columns showing the number of SNPs measured in this person (NoMeasured), the number of measured polymorphic SNPs (NoPoly), homozygosity (Hom), expected homozygosity (E(Hom)), variance, and the estimate of inbreeding, F.

Author(s)

Yurii Aulchenko, partly based on code by John Barnard

References

Purcell S. et al, (2007) PLINK: a toolset for whole genome association and population-based linkage analyses. Am. J. Hum. Genet.

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See Also

ibs, gwaa.data-class, snp.data-class

Examples

```
require(GenABEL.data)
data(ge03d2)
h <- hom(ge03d2[,c(1:100)])
h[1:5,]
homsem <- h[,"Hom"]*(1-h[,"Hom"])/h[,"NoMeasured"]
plot(h[,"Hom"],homsem)
# wrong analysis: one should use all people (for right frequency)
# and markers (for right F) available!
h <- hom(ge03d2[,c(1:10)])
h</pre>
```

hom.old

function to compute average homozygosity within a person

Description

This function computes average homozygosity (inbreeding) for a set of people, across multiple markers. Can be used for Quality Control (e.g. contamination checks)

Usage

```
hom.old(data, snpsubset, idsubset, weight="no")
```

Arguments

data Object of gwaa.data-class or snp.data-class

snpsubset Subset of SNPs to be used

idsubset People for whom average homozygosity is to be computed

weight When "no", homozygosity is computed as a proportion of homozygous geno-

types. When "freq", an estimate of inbreeding coefficint is computed (see de-

tails).

Details

With "freq" option, for person i inbreeding is estimated with

$$f_i = \frac{O_i - E_i}{(L_i - E_i)}$$

where O_i is observed homozygosity, L_i is the number of SNPs measured in individual i and

$$E_i = \sum_{j=1}^{L_i} (1 - 2p_j(1 - p_j) \frac{T_{Aj}}{T_{Aj} - 1})$$

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where T_{Aj} is the numer of measured genotypes at locus j.

Only polymorphic loci with number of measured henotypes >1 are used with this option.

This measure is the same as used by PLINK (see reference).

You should use as many people and markers as possible when estimating inbreeding from marker data.

Value

With option weight="no": A matrix with rows corresponding to the ID names and colums showing the number of genotypes measured (NoMeasured) and homozygosity (Hom).

With option weight="freq": the same as above + expected homozygosity (E(Hom)) and the estimate of inbreeding, F.

Author(s)

Yurii Aulchenko

References

Purcell S. et al, (2007) PLINK: a toolset for whole genome association and population-based linkage analyses. Am. J. Hum. Genet.

See Also

```
ibs, gwaa.data-class, snp.data-class
```

Examples

```
require(GenABEL.data)
data(ge03d2)
h <- hom(ge03d2[,c(1:100)])
homsem <- h[,"Hom"]*(1-h[,"Hom"])/h[,"NoMeasured"]
plot(h[,"Hom"],homsem)
# wrong analysis: one should use all people (for right frequency)
# and markers (for right F) available!
h <- hom(ge03d2[,c(1:10)])
h</pre>
```

HWE.show

show HWE tables

Description

This function displays HWE tables and shows Chi2 and exact HWE P-values

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Usage

```
HWE.show(data, idsubset = c(1:data@gtdata@nids),
snpsubset = c(1:data@gtdata@nsnps))
```

Arguments

data object of class gwaa.data-class or snp.data-class

snpsubset Index, character or logical vector with subset of SNPs to run analysis on. If

missing, all SNPs from data are used for analysis.

idsubset Index, character or logical vector with subset of IDs to run analysis on. If miss-

ing, all people from data/cc are used for analysis.

Value

Only screen output

Author(s)

Yurii Aulchenko

See Also

check.marker

Examples

```
require(GenABEL.data)
data(srdta)
mc <- check.marker(srdta,p.lev=0.01,ibs.mrk=0)
mc$nohwe
HWE.show(data=srdta,snps=mc$nohwe)</pre>
```

ibs

Computes (average) Idenity-by-State for a set of people and markers

Description

Given a set of SNPs, computes a matrix of average IBS for a group of individuals. This function facilitates quality control of genomic data. E.g. people with exteremly high (close to 1) IBS may indicate duplicated samples (or twins), simply high values of IBS may indicate relatives.

Usage

```
ibs(data, snpsubset, idsubset = NULL,
  cross.idsubset = NULL, weight = "no", snpfreq = NULL)
```

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Arguments

data object of snp.data-class

snpsubset Index, character or logical vector with subset of SNPs to run analysis on. If

missing, all SNPs from data are used for analysis.

idsubset IDs of people to be analysed. If missing, all people from data are used for

analysis.

cross.idsubset Parameter allowing parallel implementation. Not to be used normally. If sup-

plied together with idsubset, the ibs/kinship for all pairs between idsubset and

cross.idsubset computed.

weight "no" for direct IBS computations, "freq" to weight by allelic frequency assuming

HWE and "eVar" for empirical variance to be used

snpfreq when option weight="freq" used, you can provide fixed allele frequencies

Details

When weight "freq" is used, IBS for a pair of people i and j is computed as

$$f_{i,j} = \frac{1}{N} \sum_{k} \frac{(x_{i,k} - p_k) * (x_{j,k} - p_k)}{(p_k * (1 - p_k))}$$

where k changes from 1 to N = number of SNPs GW, $x_{i,k}$ is a genotype of ith person at the kth SNP, coded as 0, 1/2, 1 and p_k is the frequency of the "+" allele. This apparently provides an unbiased estimate of the kinship coefficient.

With "eVar" option above formula changes by using (2* empirical variance of the genotype) in the denominator. The empirical variance is computed according to the formula

$$Var(g_k) = \frac{1}{M} \Sigma_i g_{ik}^2 - E[g_k]^2$$

where M is the number of people

Only with "freq" option monomorphic SNPs are regarded as non-informative.

ibs() operation may be very lengthy for a large number of people.

Value

A (Npeople X Npeople) matrix giving average IBS (kinship) values between a pair below the diagonal and number of SNP genotype measured for both members of the pair above the diagonal.

On the diagonal, homozygosity 0.5*(1+inbreeding) is provided with option 'freq'; with option 'eVar' the diagonal is set to 0.5; the diagonal is set to homozygosity with option 'no'.

attr(computedobject,"Var") returns variance (replacing the diagonal when the object is used by egscore

Author(s)

Yurii Aulchenko

74 ibs.old

See Also

check.marker, summary.snp.data, snp.data-class

Examples

```
require(GenABEL.data)
data(ge03d2c)
set.seed(7)
# compute IBS based on a random sample of 1000 autosomal marker
selectedSnps <- sample(autosomal(ge03d2c),1000,replace=FALSE)</pre>
a <- ibs(ge03d2c,snps=selectedSnps)</pre>
a[1:5,1:5]
mds <- cmdscale(as.dist(1-a))</pre>
plot(mds)
# identify smaller cluster of outliers
km <- kmeans(mds,centers=2,nstart=1000)</pre>
cl1 <- names(which(km$cluster==1))</pre>
cl2 <- names(which(km$cluster==2))</pre>
if (length(cl1) > length(cl2)) cl1 <- cl2;</pre>
c11
# PAINT THE OUTLIERS IN RED
points(mds[cl1,],pch=19,col="red")
# compute genomic kinship matrix to be used with e.g. polygenic, mmscore, etc
a <- ibs(ge03d2c,snps=selectedSnps,weight="freq")</pre>
a[1:5,1:5]
# now replace diagonal with EIGENSTRAT-type of diaganal to be used for egscore
diag(a) <- hom(ge03d2c[,autosomal(ge03d2c)])$Var</pre>
a[1:5,1:5]
####################################
# compare 'freq' with 'eVar'
####################################
ibsFreq <- ibs(ge03d2c,snps=selectedSnps, weight="freq")</pre>
ibsEvar <- ibs(ge03d2c,snps=selectedSnps, weight="eVar")</pre>
mdsEvar <- cmdscale( as.dist( 0.5 - ibsEvar ) )</pre>
plot(mdsEvar)
outliers <- (mdsEvar[,1]>0.1)
ibsFreq[upper.tri(ibsFreq,diag=TRUE)] <- NA</pre>
ibsEvar[upper.tri(ibsEvar,diag=TRUE)] <- NA</pre>
plot(ibsEvar,ibsFreq)
points(ibsEvar[outliers,outliers],ibsFreq[outliers,outliers],col="red")
```

ibs.old

Computes (average) Idenity-by-State for a set of people and markers

Description

Given a set of SNPs, computes a matrix of average IBS for a group of people

ibs.old 75

Usage

```
ibs.old(data, snpsubset, idsubset, weight="no")
```

Arguments

data object of snp.data-class

snpsubset Index, character or logical vector with subset of SNPs to run analysis on. If

missing, all SNPs from data are used for analysis.

idsubset Index, character or logical vector with subset of IDs to run analysis on. If miss-

ing, all people from data are used for analysis.

weight "no" for direct IBS computations, "freq" to weight by allelic frequency

Details

This function facilitates quality control of genomic data. E.g. people with exteremly high (close to 1) IBS may indicate duplicated samples (or twins), simply high values of IBS may indicate relatives.

When weight "freq" is used, IBS for a pair of people i and j is computed as

$$f_{i,j} = \sum_{k} \frac{(x_{i,k} - p_k) * (x_{j,k} - p_k)}{(p_k * (1 - p_k))}$$

where k changes from 1 to N = number of SNPs GW, $x_{i,k}$ is a genotype of ith person at the kth SNP, coded as 0, 1/2, 1 and p_k is the frequency of the "+" allele. This apparently provides an unbiased estimate of the kinship coefficient.

Only with "freq" option monomorphic SNPs are regarded as non-informative.

ibs() operation may be very lengthy for a large number of people.

Value

A (Npeople X Npeople) matrix giving average IBS (kinship) values between a pair below the diagonal and number of SNP genotype measured for both members of the pair above the diagonal.

On the diagonal, homozygosity (0.5+inbreeding) is provided.

Author(s)

Yurii Aulchenko

See Also

check.marker, summary.snp.data, snp.data-class

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Examples

```
require(GenABEL.data)
data(ge03d2c)
a <- ibs(data=ge03d2c,ids=c(1:10),snps=c(1:1000))
# compute IBS based on a random sample of 1000 autosomal marker
a <- ibs(ge03d2c,snps=sample(ge03d2c@gtdata@snpnames[ge03d2c@gtdata@chromosome!="X"], a <- ibs(ge03d2c,snps=sample(ge03d2c@gtdata@snpnames[ge03d2c@gtdata@chromosome!="X"], a <- ibs(ge03d2c,snps=sample(ge03d2c@gtdata@snpnames[ge03d2c@gtdata@chromosome!="X"], a <- ibs(ge03d2c,snps=sample(ge03d2c@gtdata@snpnames[ge03d2c@gtdata@chromosome!="X"], a <- ibs(ge03d2c@gtdata@snpnames[ge03d2c@gtdata@chromosome!="X"], a <- ibs(ge03d2c@gtdata@snpnames[ge03d2c@gtdata@chromosome]="X"], a <- ibs(ge03d2c@gtdata@snpnames[ge03d2c@gtdata@snpnames[ge03d2c@gtdata@snpnames[ge03d2c@gtdata@snpnames]], a <- ibs(ge03d2c@gtdata@snpnames[ge03d2c@gtdata@snpnames]], a <- ibs(ge03d2c@gtdata@snpnames[ge03d2c@gtdata@snpnames]], a <- ibs(ge03d2c@gtdata@snpnames], a <- ibs(ge03d2c@gtdata@sn
1000, replace=FALSE), weight="freq")
mds <- cmdscale(as.dist(1-a))</pre>
plot(mds)
# identify smaller cluster of outliers
km <- kmeans(mds,centers=2,nstart=1000)</pre>
cl1 <- names(which(km$cluster==1))</pre>
cl2 <- names(which(km$cluster==2))</pre>
if (length(cl1) > length(cl2)) cl1 <- cl2;</pre>
cl1
# PAINT THE OUTLIERS IN RED
points(mds[cl1,],pch=19,col="red")
```

impute2databel

converts IMPUTE-imputed files to DatABEL (filevector) format

Description

this function converts IMPUTE-imputed files to DatABEL (filevector) format containing estimated dosages. After conversion, two files (outfile.fvi and outfile.fvd), corresponding to a single filevector object, will appear on the disk; a 'databel-class' object connected to these files will be returned to R.

Usage

```
impute2databel(genofile, samplefile, outfile,
  makeprob = TRUE, old = FALSE, dataOutType = "FLOAT")
```

Arguments

genofile IMPUTE genotype file name samplefile IMPUTE sample file name

outfile output file name

makeprob whether probability-files are also to be arranged

old for developers' use only

 ${\tt dataOutType} \qquad \quad {\tt the\ output\ data\ type,\ either\ "FLOAT"\ or\ "DOUBLE"\ (or\ another\ DatABEL/file vector\ and\ another\ DatABEL/file vector\ another\ DatABEL/file\ another\ DatABEL/f$

type)

Value

'databel-class' object

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impute2mach

converts IMPUTE to MACH files

Description

function to convert IMPUTE files to MACH format

Usage

```
impute2mach(genofile, infofile, samplefile, machbasename,
  maketextdosefile = TRUE, ...)
```

Arguments

genofile IMPUTE genotype file name infofile IMPUTE info file name samplefile IMPUTE sample file name

machbasename base name for MACH-formatted outputs

maketextdosefile

whether a text dosefile is to be generated (if not, only filevector (*.fvi / *.fvd) files, usable with ProbABEL/DatABEL, will be generated). Default: TRUE

. . . arguments passed to extract.annotation.impute (DO CHECK the documentation,

otherwise your annotation may be skrewed up!)

Value

nothing returned except files generated on the disk

Author(s)

Yurii Aulchenko

load.gwaa.data

function to load GWAA data

Description

Load data (genotypes and phenotypes) from files to gwaa.data object

Usage

```
load.gwaa.data(phenofile = "pheno.dat", genofile = "geno.raw",
force = TRUE, makemap = FALSE, sort = TRUE, id = "id")
```

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Arguments

phenofile data table with phenotypes
genofile internally formatted genotypic data file (see convert.snp.text to convert data)
force Force loading the data if heterozygous X-chromosome genotypes are found in male
makemap Make a consequtive map in case if map is provided chromosome-specifically
sort Should SNPs be sorted in ascending order according to chromosome and position?
id name of the column containing personal identification code in the phenofile

Details

The genofile must be the one resulting from convert.snp.text, convert.snp.ped, convert.snp.tped, or convert.snp.illumina (see documentation for these functions for the file formats).

The phenotype file relates study subjects with their covariate and outcome values. In the phenotypic data file, the first line gives a description of the data contained in a particular column; the names should be unique, otherwise R will change them. The first column of the phenotype file MUST contain the subjects' unique ID, named "id"; there should also be a column named "sex" and giving sex information (0 = female, 1 = male). Other columns in the file should contain phenotypic information. Missing values should be coded with "NA"; binary traits should have values 0 or 1. An example of few first lines of a phenotype file is as follows:

```
id sex age bt1 qt qt1
"289982" 0 30.33 NA NA 3.93
"325286" 0 36.514 1 0.49 3.61
"357273" 1 37.811 0 1.65 5.30
"872422" 1 20.393 0 1.95 4.07
"1005389" 1 28.21 1 0.35 3.90
```

This file tells us that, for example, person 325286 is female (0 in second column), and she has "1" (usually this means a "case") value for the trait "bt1", so on. Person 289982 has measurements only for sex, age and qt1, while other measurements are missing (NA, Not Available).

IDs are better kept in quotation (this would keep away the problem of e.g., leading zeros).

Value

Object of class gwaa.data

Author(s)

Yurii Aulchenko

See Also

save.gwaa.data, convert.snp.text, convert.snp.ped, convert.snp.tped, convert.snp.illumina

mach2databel 79

mach2databel	converts MACH-imputed files to DatABEL (filevector) format

Description

This function converts mach-imputed files to DatABEL (filevector) format. After conversion, two files (outfile.fvi and outfile.fvd), corresponding to single filevector object, will appear on the disk; 'databel-class' object connected to these files will be returned to R

Usage

```
mach2databel(imputedgenofile, mlinfofile, outfile,
  isprobfile = FALSE, dataOutType = "FLOAT")
```

Arguments

imputedgenofile

MACH mldose (or mlprob) file name

mlinfofile MACH mlinfo file name

outfile output file name

isprobfile whether imputedgenofile is a prob-file (default is FALSE, that is dose-file as-

sumed)

dataOutType the output data type, either "FLOAT" or "DOUBLE" (or other DatABEL/filevector

type)

Value

databel-class object

Author(s)

Yurii Aulchenko

makeTransitionMatrix Genotype transition probabilities matrices

Description

Function to generate genotypic transition probabilities matrices, which represent conditional probabilities P(g1lg2,nmeioses), where g1 is henotype of person one (AA, AB or BB), g2 is genotype of person two, and nmeioses is the number of meioses separating these two individuals (0 for twins, 1 for parent-offspring, c(2,2) for sibs, 2 for grandparent-grandchild pairs, etc.)

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Usage

```
makeTransitionMatrix(q, nmeioses = 1000)
```

Arguments

number of meioses separating two individuals ((a vector) of non-negative integers). If a vector, it is assumed it lists all meiotic paths connecting the pair

q (a vector of) the coded allele frequency(ies) (e.g. "Q.2" of GenABEL-package)

Value

If q is scalar, a 3x3 matrix is returned, where elements represent conditional transition probabilities P(g1|g2,nmeioses); rows correspond to the genotypes of g1, and columns correspond to the genotypes of g2. If coded allele is 'B', then e.g. element [1,2] gives the probability P(g1='AA'|g2='AB',nmeioses=nmeioses).

If q is a vector, series of above-described matrices are returned as an 'array' object. A matrix constructed for certain element q[i] can be accessed via result[,,i].

Author(s)

Yurii Aulchenko

Examples

```
# transition matrix for parent-offspring, for q=0.1
makeTransitionMatrix(0.1,nmeioses=1)
# for a set of q's
makeTransitionMatrix(c(0.1,0.9),nmeioses=1)
# for sibs
makeTransitionMatrix(0.1,nmeioses=c(2,2))
# for half-sibs (or grandparent-grandchild)
makeTransitionMatrix(0.1,nmeioses=2)
# for remote relatives
makeTransitionMatrix(0.1,nmeioses=10)
# for independent
makeTransitionMatrix(0.1,nmeioses=1000)
```

merge.gwaa.data

function to merge objects of gwaa.data-class

Description

function to merge two objects of gwaa.data-class

Usage

```
## S3 method for class 'gwaa.data' merge(x, y, ...)
```

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Arguments

```
x the first object of gwaa.data-classy the second object of gwaa.data-class... arguments passed to merge.snp.data
```

Details

This function calls merge.snp.data to merge gtdata slots of the incoming objects; the phenotypic data contained in phdata slots are merged using standard function for data frames with arguments by="id" and all=TRUE. The merged object is filtered and sorted according to order of ids presented in the merged snp.data object.

Value

```
Merged object of gwaa.data-class
```

Author(s)

Maksim Struchalin, Yurii Aulchenko

See Also

```
merge.snp.data,add.phdata
```

Examples

```
require(GenABEL.data)
data(srdta)
x1 <- srdta[c(1,3,5,6),c(2,4,5,6)]
x2 <- srdta[c(2,4,5,6),c(1,3,5,6)]
x3 <- merge(x1,x2)
x1
as.character(x1)
x2
as.character(x2)
x3
as.character(x3)
srdta[1:6,1:6]
as.character(srdta[1:6,1:6])</pre>
```

merge.snp.data

function to merge objects of snp.data-class

Description

function to merge two objects of snp.data-class

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Usage

```
## S3 method for class 'snp.data'
merge(x, y, ..., error_amount = 1e+06, replacena = TRUE,
forcestranduse = FALSE, sort = TRUE,
intersected_snps_only = FALSE)
```

Arguments

x the first object of snp.data-class
y the second object of snp.data-class
... additional arguments (not used or passed)
error_amount if this amount of errors is exceeded, only error table is returned
replacena Some genotypes may be missing in set 1, but measured in set 2. If replacena=TRUE, genotypes from the set 2 will appear in the merged data.

forcestranduse if TRUE, forces use of strand infomation even if coding information is sufficient for merging
sort if TRUE, sorts the object according to chromosome and SNP position intersected_snps_only

if TRUE, then only interesected SNPs will be in output

Details

By default, when a genotype for a person is measured in both set "x" and set "y", and these are not equal, the value specified by set "x" is returned in the merged set. In case when genotype is NA in the first set, the behaviour depends on the value of the "replacena" parameter – if set to TRUE (default), these are replaced with the non-NA values from set "y".

When "forcestranduse" is set to FALSE, strand information is not used unless the coding is not sufficient for merging the data (i.e. starnd information is used only to merge A/T and G/C polymorphisms).

SNP error is returned to "snp" table when SNP coding is incompatible between the two sets. For such SNPs, only the data provided by set "x" are used in the merged data.

ID error is returned to "id" table when genotypes of the same person at the same SNP are different between set "x" and set "y". For such genotypes, the data provided by set "x" are used in the merged data.

Value

A list is returned

id

This table summarises individual genotype inconsistencies. These may occur when some person is present and genotyped for the same marker in both sets, but these genotypes are inconsistent. The table's first column, "id", contains personal ID, the second, "snpnames", contain SNP name, and third (set "x") and fourth (set "y") contain the genotypes for this person at this SNP in sets 1 and 2.

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snp This table summarises coding errors. These occur when for some SNPs coding

in the set 1 is not compatible with set 2. The table's first column ("snpnames") provides SNP name, and second (set "x") and third (set "y") report coding used

in respective sets.

data merged object of snp.data-class

Author(s)

Maksim Struchalin, Yurii Aulchenko

See Also

```
merge.gwaa.data, add.phdata
```

Examples

```
require(GenABEL.data)
data(srdta)
x1 <- srdta[c(1,3,5,6),c(2,4,5,6)]@gtdata
x2 <- srdta[c(2,4,5,6),c(1,3,5,6)]@gtdata
x3 <- merge(x1,x2)
as.character(x1)
as.character(x2)
as.character(x3$data)
as.character(srdta[1:6,1:6])</pre>
```

mlreg

Linear and logistic regression and Cox models for genome-wide SNP data

Description

Linear and logistic regression and Cox models for genome-wide SNP data

Usage

```
mlreg(formula, data, gtmode = "additive", trait.type = "guess", propPs = 1)
```

Arguments

formula Standard formula object

data an object of gwaa.data-class

gtmode Either "additive", "dominant", "recessive" or "overdominant". Specifies the

analysis model.

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trait.type Either "gaussian", "binomial" or "survival", corresponding to analysis using lin-

ear regression, logistic regression, and Cox proportional hazards models, respectively. When default vale "guess" is used, the program tries to guess the

type

propPs proportion of non-corrected P-values used to estimate the inflation factor Lambda,

passed directly to the estlambda

Details

Linear regression is performed using standard approach; logisite regression is implemented using IRLS; Cox model makes use of code contributed by Thomas Lumley (survival package).

For logistic and Cox, exp(effB) gives Odds Ratios and Hazard Ratios, respectively.

Value

```
An object of scan.gwaa-class
```

Author(s)

Yurii Aulchenko

See Also

```
GASurv, qtscore
```

Examples

```
require(GenABEL.data)
data(ge03d2)
dta <- ge03d2[,1:100]
# analysis using linear model
xq <- mlreg(bmi~sex,dta)
# logistic regression, type guessed automatically
xb <- mlreg(dm2~sex,dta)
# Cox proportional hazards model, assuming that age is the follow-up time
# generally this does not make sense (could be ok if age is age at onset)
xs <- mlreg(GASurv(age,dm2)~sex,dta)</pre>
```

mlreg.p

EXPERIMENTAL Linear and logistic regression and Cox models for genome-wide SNP data

Description

Linear and logistic regression and Cox models for genome-wide SNP data

mlreg.p

Usage

```
mlreg.p(formula, data, snpsubset, idsubset, gtmode = "additive", trait.type = "guess")
```

Arguments

formula Standard formula object data an object of gwaa.data-class snpsubset Index, character or logical vector with subset of SNPs to run analysis on. If missing, all SNPs from data are used for analysis. idsubset Index, character or logical vector with subset of IDs to run analysis on. If missing, all people from data/cc are used for analysis. Either "additive", "dominant", "recessive" or "overdominant". Specifies the gtmode analysis model. Either "gaussian", "binomial" or "survival", corresponding to analysis using lintrait.type ear regression, logistic regression, and Cox proportional hazards models, respectively. When default vale "guess" is used, the program tries to guess the

Details

Linear regression is performed using standard approach; logisite regression is implemented using IRLS; Cox model makes use of code contributed by Thomas Lumley (survival package).

For logistic and Cox, exp(effB) gives Odds Ratios and Hazard Ratios, respectively.

Value

An object of scan.gwaa-class

type

Author(s)

Yurii Aulchenko

See Also

GASurv, qtscore

Examples

```
require(GenABEL.data)
data(ge03d2)
dta <- ge03d2[,1:100]
# analysis using linear model
xq <- mlreg.p(bmi~sex,dta)
# logistic regression, type guessed automatically
xb <- mlreg.p(dm2~sex,dta)
# Cox proportional hazards model, assuming that age is the follow-up time
# generally this does not make sense (could be ok if age is age at onset)
xs <- mlreg.p(GASurv(age,dm2)~sex,dta)</pre>
```

86 mmscore

Description

Score test for association between a trait and genetic polymorphism, in samples of related individuals

Usage

```
mmscore(h2object,data,snpsubset,idsubset,strata,times=1,quiet=FALSE,
bcast=10,clambda=TRUE,propPs=1.0)
```

Arguments

h2object	An object returned by polygenic polygenic mixed model analysis routine. The sub-objects used are measuredIDs, residualY, and InvSigma. One can supply mmscore with a fake h2object, containing these list elements.
data	An object of gwaa.data-class. ALWAYS PASS THE SAME OBJECT WHICH WAS USED FOR ipolygenic ANALYSIS, NO SUB-SETTING IN IDs (USE IDSUBSET ARGUMENT FOR SUB-SETTING)!!!
snpsubset	Index, character or logical vector with subset of SNPs to run analysis on. If missing, all SNPs from data are used for analysis.
idsubset	Index, character or logical vector with subset of IDs to run analysis on. If missing, all people from data/cc are used for analysis.
strata	Stratification variable. If provieded, scores are computed within strata and then added up.
times	If more then one, the number of replicas to be used in derivation of empirical genome-wide significance. NOTE: The structure of the data is not exchangable, therefore do not use times > 1 unless you are really sure you understand what you are doing!
quiet	do not print warning messages
bcast	If the argument times > 1, progress is reported once in bcast replicas
clambda	If inflation facot Lambda is estimated as lower then one, this parameter controls if the original P1df (clambda=TRUE) to be reported in Pc1df, or the original 1df statistics is to be multiplied onto this "deflation" factor (clambda=FALSE). If a numeric value is provided, it is used as a correction factor.
propPs	proportion of non-corrected P-values used to estimate the inflation factor Lambda, passed directly to the ${\tt estlambda}$

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Details

Score test is performed using the formula

$$\frac{((G-E[G])V^{-1}residualY)^2}{(G-E[G])V^{-1}(G-E[G])}$$

where G is the vector of genotypes (coded 0, 1, 2) and E[G] is a vector of (strata-specific) mean genotypic values; V^{-1} is the InvSigma and residualY are residuals from the trait analysis with polygenic procedure.

This test is similar to that implemented by Abecasis et al. (see reference).

Value

Object of class scan. gwaa-class; only 1 d.f. test is implemented currently.

Author(s)

Yurii Aulchenko

References

Chen WM, Abecasis GR. Family-based association tests for genome-wide association scans. Am J Hum Genet. 2007 Nov;81(5):913-26.

See Also

```
grammar, qtscore, egscore, plot.scan.gwaa, scan.gwaa-class
```

Examples

```
# ge03d2 is rather bad data set to demonstrate,
# because this is a population-based study
require(GenABEL.data)
data(ge03d2.clean)
#take half for speed
ge03d2.clean <- ge03d2.clean[1:100,]</pre>
gkin <- ibs(ge03d2.clean,w="freq")</pre>
h2ht <- polygenic(height ~ sex + age,kin=gkin,ge03d2.clean)
h2ht$est
mm <- mmscore(h2ht,data=ge03d2.clean)
# compute grammar
gr <- qtscore(h2ht$pgres,data=ge03d2.clean,clam=FALSE)</pre>
gc <- qtscore(height ~ sex + age,data=ge03d2.clean)</pre>
#compare
plot(mm,df="Pc1df",cex=0.5)
add.plot(gc,df="Pc1df",col="red")
add.plot(gr,df="Pc1df",col="lightgreen",cex=1.1)
# can see that mmscore and grammar are quite the same... in contrast to GC
```

88 npsubtreated

npsubtreated	non-parametric trait "imputations" in treated people
	non puramente trans imputantens in medica people

Description

For people on treatment, the algorithm substitutes the value of the trait using non-parametric algorithm described in Tobin et al., 2005. This algorithm assumes that the measurement in treated subject is a right-censored trait. Essentially, the algorithm substitutes the QT for a person on treatment with the mean of the above-ranked substituted QT value.

Usage

```
npsubtreated(trait, medication, increase = FALSE)
```

Arguments

trait vector if trait values

medication medication indicator (0/1)

increase if medication INCREASE the value of the trait (should never be true for e.g.

blood pressure, LDL cholesterol, etc.)

Details

Should put the formulas here...

Value

Vector of trait walues, where the values for treated people are substituted with average of the above-ranked substituted trait value.

Author(s)

Yurii Aulchenko

References

Levy D, DeStefano AL, Larson MG, O'Donnell CJ, Lifton RP, Gavras H, Cupples LA, Myers RH. Evidence for a gene influencing blood pressure on chromosome 17. Genome scan linkage results for longitudinal blood pressure phenotypes in subjects from the framingham heart study. Hypertension. 2000 Oct;36(4):477-83.

Tobin MD, Sheehan NA, Scurrah KJ, Burton PR. Adjusting for treatment effects in studies of quantitative traits: antihypertensive therapy and systolic blood pressure. Stat Med. 2005 Oct 15;24(19):2911-35.

patch_strand 89

Examples

```
# simulate SBP data
simmeddat <- function(mu=144,bage=0.5,bsex=4.,bg=2.,pB=0.3,rvar=21^2,N=1000) {
ageb <- c(25,74)
pmale < - .5
htthresh <- 160
trprob <- .5
mutreff <- (-15.)
trvar <- 4^2
age <- runif(N,min=ageb[1],max=ageb[2])</pre>
sex <- 1*(runif(N)<=pmale)</pre>
gt <- rbinom(N,size=2,prob=pB)</pre>
y.true <- rnorm(N,mu,sqrt(rvar)) + bage*age + bsex*sex + bg*gt</pre>
d.true <- (y.true>=htthresh)
medication <- 1*d.true
medication[d.true] <- 1*(runif(sum(d.true))<=trprob)</pre>
treatm <- rnorm(sum(medication), mutreff, sqrt(trvar))</pre>
treatm[treatm>0] <- 0</pre>
treff <- rep(0,N)</pre>
treff[medication==1] <- treatm</pre>
y.obs <- y.true + treff
out <- data.frame(age,sex,gt,y.true,d.true,medication,treff,y.obs)</pre>
}
x <- simmeddat(bg=2.0,N=3000)</pre>
x[1:15,]
# substitute value of treated people
imptra <- npsubtreated(x$y.obs,x$medication)</pre>
imptra[1:15]
# Almost always, correlation should be higher for the "imputed" trait
cor(x$y.true,x$y.obs)
cor(x$y.true,imptra)
# see what comes out of regression
# analysis of true value
summary(lm(y.true~sex+age+gt,data=x))
# ignore treatment (as a rule, betas are underestimated;
# on average, power goes down)
summary(lm(y.obs~sex+age+gt,data=x))
# treatment as covariate (as a rule, betas are underestimated;
# on average, power goes down)
summary(lm(y.obs~sex+age+gt+medication,data=x))
# analyse "imputed" trait (as a rule betas are better; power
# approaches that of analysis of "true" trait)
summary(lm(imptra~sex+age+gt,data=x))
```

90 perid.summary

Description

Changes strand in gwaa.data-class object

Usage

```
patch_strand(data,snpid,strand,based_on="snpnames", quiet = TRUE)
```

Arguments

data gwaa.data or snp.data object

snpid vector of ids of snsp (name or position)

strand vector of strands ("+","-","u")

based_on either "snpnames" or "map" depending on what info is provided by snpid

quiet indicates if recoding report should be directed to the screen

Details

For SNPs, as identified by 'snpid', changes strand to strand specified by 'strand'

Value

object of gwaa.data or snp.data class

Author(s)

Yurii Aulchenko

Examples

```
require(GenABEL.data)
data(srdta)
as.character(srdta@gtdata@strand[1:20])
a <- patch_strand(srdta,srdta@gtdata@snpnames[1:10],rep("+",10))
as.character(a@gtdata@strand[1:20])
a <- patch_strand(srdta,srdta@gtdata@map[1:10],rep("+",10),based_on="map")
as.character(a@gtdata@strand[1:20])</pre>
```

perid.summary

Summary of marker data per person

Description

Produces call rate and heterozygosity per person

Usage

```
perid.summary(data, snpsubset, idsubset, ...)
```

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Arguments

data	object of snp.data-class
snpsubset	Index, character or logical vector with subset of SNPs to run analysis on. If missing, all SNPs from data are used for analysis.
idsubset	Index, character or logical vector with subset of IDs to run analysis on. If missing, all people from data are used for analysis.
	additional parameters to be passed to hom

Details

This function facilitates quality control of genomic data. E.g. extreme outliers for heterozygosity indicate possibly contaminated DNA samples, while low call rate of a person may indicate poor DNA quality.

Value

A matrix, giving per person (row) its' average heterozygosity ("Het" column) and call rate ("CallPP"), over all SNPs

Author(s)

Yurii Aulchenko

See Also

```
check.marker, summary.snp.data, snp.data-class
```

Examples

```
require(GenABEL.data)
data(ge03d2c)
a <- perid.summary(data=ge03d2c,snps=c(1:100),ids=c(1:10))
a
a <- perid.summary(data=ge03d2c)
hist(a[,"CallPP"])
hist(a[,"Het"])</pre>
```

PGC Polynomial genomic control

Description

This function estimates the genomic controls for different models and degrees of freedom, using polinomial function. Polinomial coefficients are estimated by optimizing different error functions: regress, median, ks.test or group regress.

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Usage

```
PGC(data, method = "group_regress", p, df, pol.d = 3,
  plot = TRUE, index.filter = NULL, start.corr = FALSE,
  proportion = 1, n_quiantile = 5, title_name = "Lambda",
  type_of_plot = "plot", lmax = NULL, color = "red")
```

Arguments

data Input vector of Chi square statistic method Function of error to be optimized. Can be "regress", "median", "ks.test" or "group_regress" Input vector of allele frequencies р df Number of degrees of freedom The degree of polinomial function pol.d plot If TRUE, plot of lambda will be produced For regress method use it only when you want to make calculations faster start.corr index.filter Index of variables in data vector, that will be used in analysis if zero - all variables will be used The proportion of lowest P (Chi2) to be used when estimating the inflation factor proportion Lambda for "regress" method only The number of groups for "group_regress" method n_quiantile The title name for plot title_name type_of_plot For developers only The threshold for lambda for plotting (optional) lmax

Value

color

A list with elements

data Output vector corrected Chi square statistic

The color of the plot

b Polinomial coefficients

Author(s)

Yakov Tsepilov

Examples

```
require(GenABEL.data)
data(ge03d2)
ge03d2 <- ge03d2[seq(from=1,to=nids(ge03d2),by=2),seq(from=1,to=nsnps(ge03d2),by=3)]
qts <- mlreg(dm2~1,data=ge03d2,gtmode = "additive")
chi2.1df <- results(qts)$chi2.1df
s <- summary(ge03d2)
freq <- s$0.2
result=PGC(data=chi2.1df,method="median",p=freq,df=1, pol.d=2, plot=TRUE, lmax=1.1,start.corr=FALSE)</pre>
```

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plot.check.marker

plots "check.marker" object

Description

Plots "check.marker" object, as returned by check.marker

Usage

```
## S3 method for class 'check.marker'
plot(x, y, ...)
```

Arguments

x Object of class "check.marker", as returned by check.marker or snp. subset

y this argument is not used

... other arguments to be passed to plot

Details

In this plot, along the X axes, you can see colour representation of markers which did not pass (pass – black) the QC. The diagonal shows redundant markers. If for some marker there exist markers, which show exactly the same (or some minimum concordance) genotypic distribution, such markers are depicted as crosses an solid line is dropped on the X axes from it. Other solid line connects the original SNP with the redundant ones (depicted as circles). From each redundant SNP, a dashed line is dropped on X. Normally, one expects that redundant markers are positioned very closely and redundancy appears because of linkage disequilibrium.

Value

No value returned. Explanatory note is shown on the screen.

Author(s)

Yurii Aulchenko

See Also

```
check.marker, snp.subset
```

Examples

```
require(GenABEL.data)
data(srdta)
mc <- check.marker(data=srdta@gtdata[,1:100],redundant="all",maf=0.01,
minconcordance=0.9,fdr=.1,ibs.mrk=0)
mc <- check.marker(data=srdta@gtdata[,1:100],maf=0.01,fdr=.1,ibs.mrk=0)
plot(mc)</pre>
```

94 plot.scan.gwaa

```
mc1 <- snp.subset(mc,snps=srdta@gtdata@snpnames[20:40])
plot(mc1)</pre>
```

plot.scan.gwaa

function to plot GWAA results

Description

Plots results of GWA analysis

Usage

```
## S3 method for class 'scan.gwaa'
plot(x, y, ..., df=1, ystart=0, col=c("blue","green"),
sort=TRUE, ylim, delta = 1 , main = getcall(x))
```

Arguments

X	object of type scan.gwaa-class, as returned by scan.glm, qtscore, ccfast, emp.ccfast, emp.qtscore, or scan.haplo
у	this argument is not used
	additional arguments to be passed to plot
df	Plot results of 1 or 2-df test (1, 2). Could be also "Pc1df" (for GC corrected P-values) and "Pc2df" (for robust genomic control of the 2 df test)
ystart	truncate lower value of Y at this point (can help avoiding plotting too may points)
ylim	ylim, same as in the standard plot function
col	which colors to use to depict consecutive chromosomes
sort	whether results should be plotted after sorting by chromosome and position
delta	gap width between chromosomes
main	title of the plot

Value

No value returned.

Author(s)

Yurii Aulchenko

See Also

```
\verb|scan.gwaa-class|, \verb|add.plot|, \verb|snp.subset|, \verb|scan.glm|, \verb|qtscore|, \verb|ccfast|, emp.qtscore|, emp.ccfast|, \verb|scan.haplo||
```

plot.scan.gwaa.2D 95

Examples

```
require(GenABEL.data)
data(srdta)
a <- qtscore(bt,srdta,snps=c(1:250))
plot(a)
add.plot(a,df="Pc1df",col="green")</pre>
```

plot.scan.gwaa.2D

function to plot 2D scan results

Description

Plots results of 2D analysis produced by scan.glm.2D or scan.haplo.2D

Usage

```
## S3 method for class 'scan.gwaa.2D'
plot(x, y, ..., df=1)
```

Arguments

x	object of type scan.gwaa.2D-class, as returned by scan.glm.2D or scan.haplo.2D
у	this argument is not used
	additional arguments to be passed to plot
df	Whether 1, 2, or "all" d.f.s should be plotted. Note that for scan.haplo.2D 1 and 2 d.f. list the same values.

Details

Now plots only "allelelic" results. This is fine for scan.haplo.2Das only alleic tests are produced; however, scan.glm.2D also produces "genotypic" results.

Value

No value returned.

Author(s)

Yurii Aulchenko

See Also

```
scan.gwaa.2D-class, scan.glm.2D, scan.haplo.2D
```

Examples

```
require(GenABEL.data)
data(srdta)
a <- scan.glm.2D("qt3~CRSNP",data=srdta,snps=c(1:10))
# "allelic" results
plot(a)
# to plot "genotypic" results:
filled.contour(x=a$map,y=a$map,z=-log10(a$P2df))</pre>
```

polygenic

Estimation of polygenic model

Description

This function maximises the likelihood of the data under polygenic model with covariates an reports twice negative maximum likelihood estimates and the inverse of the variance-covariance matrix at the point of ML.

Usage

```
polygenic(formula, kinship.matrix, data, fixh2,
  starth2 = 0.3, trait.type = "gaussian",
  opt.method = "nlm", scaleh2 = 1, quiet = FALSE,
  steptol = 1e-08, gradtol = 1e-08, optimbou = 8,
  fglschecks = TRUE, maxnfgls = 8, maxdiffgls = 1e-04,
  patchBasedOnFGLS = TRUE, llfun = "polylik_eigen",
  eigenOfRel, ...)
```

Arguments

formula	Formula describing fixed effects to be used in the analysis, e.g. $y \sim a + b$ means that outcome (y) depends on two covariates, a and b. If no covariates used in the analysis, skip the right-hand side of the equation.
kinship.matrix	Kinship matrix, as provided by e.g. ibs(,weight="freq"), or estimated outside of GenABEL from pedigree data.
data	An (optional) object of gwaa.data-class or a data frame with outcome and covariates
fixh2	Optional value of heritability to be used, instead of maximisation. The uses of this option are two-fold: (a) testing significance of heritability and (b) using a priori known heritability to derive the rest of MLEs and varcov. matrix.
starth2	Starting value for h2 estimate
trait.type	"gaussian" or "binomial"
opt.method	"nlm" or "optim". These two use different optimisation functions. We suggest using the default nlm, although optim may give better results in some situations

scaleh2 Only relevant when "nlm" optimisation function is used. "scaleh2" is the her-

itability scaling parameter, regulating how "big" are parameter changes in h2 with respect to changes in other parameters. As other parameters are estimated from previous regression, these are expected to change little from the initial estimate. The default value of 1000 proved to work rather well under a range of

conditions.

quiet If FALSE (default), details of optimisation process are reported

steptol steptal parameter of "nlm" gradtol gradtol parameter of "nlm"

optimbou fixed effects boundary scale parameter for 'optim'

fglschecks additional check for convergence on/off (convergence between estimates ob-

tained and that from FGLS)

maxnfgls number of fgls checks to perform
maxdiffgls max difference allowed in fgls checks

patchBasedOnFGLS

if FGLS checks not passed, 'patch' fixed effect estimates based on FGLS expec-

tation

11fun function to compute likelihood (default 'polylik_eigen', also available – but not

recommended – 'polylik')

eigenOfRel results of eigen(relationship matrix = 2*kinship.matrix). Passing this can de-

crease computational time substantially if multiple traits are analysed using the same kinship matrix. This option will not work if any NA's are found in the trait and/or covariates and if the dimensions of the 'eigen'-object, trait, covariates,

kinship do not match.

... Optional arguments to be passed to nlm or (optim) minimisation function

Details

One of the major uses of this function is to estimate residuals of the trait and the inverse of the variance-covariance matrix for further use in analysis with mmscore and grammar.

Also, it can be used for a variant of GRAMMAR analysis, which allows for permutations for GW significance by use of environmental residuals as an analysis trait with qtscore.

"Environmental residuals" (not to be mistaken with just "residuals") are the residual where both the effect of covariates AND the estimated polygenic effect (breeding values) are factored out. This thus provides an estimate of the trait value contributed by environment (or, turning this other way around, the part of the trait not explained by covariates and by the polygene). Polygenic residuals are estimated as

$$\sigma^2 V^{-1} (Y - (\hat{\mu} + \hat{\beta} C_1 + ...))$$

where $sigma^2$ is the residual variance, V^{-1} is the InvSigma (inverse of the var-cov matrix at the maximum of polygenic model) and $(Y - (\hat{\mu} + \hat{\beta}C_1 + ...))$ is the trait values adjusted for covariates (also at at the maximum of polygenic model likelihood).

It can also be used for heritability analysis. If you want to test significance of heritability, estimate the model and write down the function minimum reported at the "h2an" element of the output (this

is twice the negative MaxLikelihood). Then do a next round of estimation, but set fixh2=0. The difference between your function minima gives a test distributed as chi-squared with 1 d.f.

The way to compute the likelihood is partly based on the paper of Thompson (see refs), namely instead of taking the inverse of the var-cov matrix every time, eigenvectors of the inverse of G (taken only once) are used.

Value

A list with values

h2an	A list supplied by the nlm minimisation routine. Of particular interest are elements "estimate" containing parameter maximal likelihood estimates (MLEs) (order: mean, betas for covariates, heritability, (polygenic + residual variance)). The value of twice negative maximum log-likelihood is returned as h2an\\$minimum.
esth2	Estimate (or fixed value) of heritability
residualY	Residuals from analysis, based on covariate effects only; NOTE: these are NOT grammar "environmental residuals"!
pgresidualY	Environmental residuals from analysis, based on covariate effects and predicted breeding value.
grresidualY	GRAMMAR+ transformed trait residuals
grammarGamma	list with GRAMMAR-gamma correction factors
InvSigma	Inverse of the variance-covariance matrix, computed at the MLEs – these are used in mmscore and grammar functions.
call	The details of call
measuredIDs	Logical values for IDs who were used in analysis (traits and all covariates measured) == TRUE
convFGLS	was convergence achieved according to FGLS criterionE

Note

Presence of twins may complicate your analysis. Check the kinship matrix for singularities, or rather use check.marker for identification of twin samples. Take special care in interpretation.

If a trait (no covariates) is used, make sure that the order of IDs in the kinship.matrix is exactly the same as in the outcome

Please note that there is alternative to 'polygenic', polygenic_hglm, which is faster than polygenic() with the llfun='polylik' option, but slightly slower than the default polygenic().

Author(s)

Yurii Aulchenko, Gulnara Svischeva

References

Thompson EA, Shaw RG (1990) Pedigree analysis for quantitative traits: variance components without matrix inversion. Biometrics 46, 399-413.

for original GRAMMAR

Aulchenko YS, de Koning DJ, Haley C. Genomewide rapid association using mixed model and regression: a fast and simple method for genome-wide pedigree-based quantitative trait loci association analysis. Genetics. 2007 177(1):577-85.

for GRAMMAR-GC

Amin N, van Duijn CM, Aulchenko YS. A genomic background based method for association analysis in related individuals. PLoS ONE. 2007 Dec 5;2(12):e1274.

for GRAMMAR-Gamma

Svischeva G, Axenovich TI, Belonogova NM, van Duijn CM, Aulchenko YS. Rapid variance components-based method for whole-genome association analysis. Nature Genetics. 2012 44:1166-1170. doi:10.1038/ng.2410

for GRAMMAR+ transformation

Belonogova NM, Svishcheva GR, van Duijn CM, Aulchenko YS, Axenovich TI (2013) Region-Based Association Analysis of Human Quantitative Traits in Related Individuals. PLoS ONE 8(6): e65395. doi:10.1371/journal.pone.0065395

See Also

```
polygenic_hglm, mmscore, grammar
```

Examples

```
# note that procedure runs on CLEAN data
require(GenABEL.data)
data(ge03d2ex.clean)
gkin <- ibs(ge03d2ex.clean,w="freq")</pre>
h2ht <- polygenic(height ~ sex + age, kin=gkin, ge03d2ex.clean)
# estimate of heritability
h2ht$esth2
# other parameters
h2ht$h2an
# the minimum twice negative log-likelihood
h2ht$h2an$minimum
# twice maximum log-likelihood
-h2ht$h2an$minimum
# for binary trait (experimental)
h2dm <- polygenic(dm2 ~ sex + age, kin=gkin, ge03d2ex.clean, trait="binomial")
# estimated parameters
h2dm$h2an
```

100 polygenic_hglm

Description

Estimation of polygenic model using a hierarchical generalized linear model (HGLM; Lee and Nelder 1996. hglm package; Ronnegard et al. 2010).

Usage

```
polygenic_hglm(formula, kinship.matrix, data,
  family = gaussian(), conv = 1e-06, maxit = 100, ...)
```

Arguments

formula	Formula describing fixed effects to be used in analysis, e.g. $y \sim a + b$ means that outcome (y) depends on two covariates, a and b. If no covariates used in analysis, skip the right-hand side of the equation.
kinship.matrix	Kinship matrix, as provided by e.g. ibs(,weight="freq"), or estimated outside of GenABEL from pedigree data.
data	An (optional) object of $gwaa.data\text{-}class$ or a data frame with outcome and covariates.
family	a description of the error distribution and link function to be used in the mean part of the model (see family for details of family functions).
conv	'conv' parameter of hglm (stricter than default, for great precision, use 1e-8).
maxit	'maxit' parameter of hglm (stricter than default).
	other parameters passed to hglm call

Details

The algorithm gives extended quasi-likelihood (EQL) estimates for restricted maximum likelihood (REML) (Ronnegard et al. 2010). Implemented is the inter-connected GLM interpretation of HGLM (Lee and Nelder 2001). Testing on fixed and random effects estimates are directly done using inter-connected GLMs. Testing on dispersion parameters (variance components) can be done by extracting the profile likelihood function value of REML.

Author(s)

Xia Shen, Yurii Aulchenko

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References

Ronnegard, L, Shen, X and Alam, M (2010). hglm: A Package For Fitting Hierarchical Generalized Linear Models. *The R Journal*, **2**(2), 20-28.

Lee, Y and Nelder JA (2001). Hierarchical generalised linear models: A synthesis of generalised linear models, random-effect models and structured dispersions. *Biometrika*, **88**(4), 987-1006.

Lee, Y and Nelder JA (1996). Hierarchical Generalized Linear Models. *Journal of the Royal Statistical Society. Series B*, **58**(4), 619-678.

See Also

polygenic, mmscore, grammar

Examples

```
require(GenABEL.data)
data(ge03d2ex.clean)
set.seed(1)
df <- ge03d2ex.clean[,sample(autosomal(ge03d2ex.clean),1000)]</pre>
gkin <- ibs(df,w="freq")</pre>
# ---- for quantitative traits
h2ht <- polygenic_hglm(height ~ sex + age, kin = gkin, df)
# ---- estimate of heritability
h2ht$esth2
# ---- other parameters
h2ht$h2an
# ---- test the significance of fixed effects
# ---- (e.g. quantitative trait)
h2ht <- polygenic_hglm(height ~ sex + age, kin = gkin, df)
# ---- summary with standard errors and p-values
summary(h2ht$hglm)
# ---- output for the fixed effects part
# ...
# MEAN MODEL
# Summary of the fixed effects estimates
             Estimate Std. Error t value Pr(>|t|)
# (Intercept) 169.53525 2.57624 65.807 < 2e-16 ***
# sex 14.10694 1.41163 9.993 4.8e-15 ***
             # age
# Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# Note: P-values are based on 69 degrees of freedom
# ---- extract fixed effects estimates and standard errors
h2ht$h2an
# ---- test the significance of polygenic effect
nullht <- lm(height ~ sex + age, df)</pre>
11 <- h2ht$ProfLogLik</pre>
10 <- as.numeric(logLik(nullht))</pre>
```

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```
# the likelihood ratio test (LRT) statistic
(S <- -2*(10 - 11))
# 5% right-tailed significance cutoff
# for 50:50 mixture distribution between point mass 0 and \chi^{2}(1)
# Self, SG and Liang KY (1987) Journal of the American Statistical Association.
qchisq(((1 - .05) - .50)/.50, 1)
# p-value
pchisq(S, 1, lower.tail = FALSE)/2
# ---- for binary traits
h2dm <- polygenic_hglm(dm2 ~ sex + age, kin = gkin, df, family = binomial(link = 'logit'))
# ---- estimated parameters
h2dm$h2an</pre>
```

qtscore

Fast score test for association

Description

Fast score test for association between a trait and genetic polymorphism

Usage

```
qtscore(formula, data, snpsubset, idsubset, strata,
  trait.type = "gaussian", times = 1, quiet = FALSE,
  bcast = 10, clambda = TRUE, propPs = 1, details = TRUE)
```

Arguments

formula	Formula describing fixed effects to be used in analysis, e.g. $y \sim a + b$ means that outcome (y) depends on two covariates, a and b. If no covariates used in analysis, skip the right-hand side of the equation.
data	An object of gwaa.data-class
snpsubset	ndex, character or logical vector with subset of SNPs to run analysis on. If missing, all SNPs from data are used for analysis.
idsubset	ndex, character or logical vector with subset of IDs to run analysis on. If missing, all people from data/cc are used for analysis.
strata	Stratification variable. If provieded, scores are computed within strata and then added up.
trait.type	"gaussian" or "binomial" or "guess" (later option guesses trait type)
times	If more than one, the number of replicas to be used in derivation of empirical genome-wide significance. See <pre>emp.qtscore</pre> , which calls qtscore with times>1 for details
quiet	do not print warning messages
bcast	If the argument times > 1, progress is reported once in bcast replicas

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clambda If inflation facot Lambda is estimated as lower then one, this parameter controls if the original P1df (clambda=TRUE) to be reported in Pc1df, or the original 1df statistics is to be multiplied onto this "deflation" factor (clambda=FALSE). If a numeric value is provided, it is used as a correction factor.

propPs proportion of non-corrected P-values used to estimate the inflation factor Lambda,

passed directly to the estlambda

when FALSE, SNP and ID names are not reported in the returned object (saves some memory). This is experimental and will be not mantained anymore as

soon as we achieve better memory efficiency for storage of SNP and ID names

(currently default R character data type used)

Details

details

When formula contains covariates, the traits is analysed using GLM and later residuals used when score test is computed for each of the SNPs in analysis. Coefficients of regression are reported for the quantitative trait.

For binary traits, odds ratios (ORs) are reportted. When adjustemnt is performed, first, "response" residuals are estimated after adjustment for covariates and scaled to [0,1]. Reported effects are approximately equal to ORs expected in logistic regression model.

With no adjustment for binary traits, 1 d.f., the test is equivalent to the Armitage test.

This is a valid function to analyse GWA data, including X chromosome. For X chromosome, stratified analysis is performed (strata=sex).

Value

Object of class scan.gwaa-class

Author(s)

Yurii Aulchenko

References

Aulchenko YS, de Koning DJ, Haley C. Genomewide rapid association using mixed model and regression: a fast and simple method for genome-wide pedigree-based quantitative trait loci association analysis. Genetics. 2007 177(1):577-85.

Amin N, van Duijn CM, Aulchenko YS. A genomic background based method for association analysis in related individuals. PLoS ONE. 2007 Dec 5;2(12):e1274.

See Also

mlreg, mmscore, egscore, emp.qtscore, plot.scan.gwaa, scan.gwaa-class

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Examples

```
require(GenABEL.data)
data(srdta)
#qtscore with stratification
a <- qtscore(qt3~sex,data=srdta)
plot(a)
b <- qtscore(qt3,strata=phdata(srdta)$sex,data=srdta)</pre>
add.plot(b,col="green",cex=2)
# qtscore with extra adjustment
a <- qtscore(qt3~sex+age,data=srdta)
plot(a)
# compare results of score and chi-square test for binary trait
a1 <- ccfast("bt",data=srdta,snps=c(1:100))</pre>
a2 <- qtscore(bt,data=srdta,snps=c(1:100),trait.type="binomial")</pre>
plot(a1, ylim=c(0, 2))
add.plot(a2,col="red",cex=1.5)
# the good thing about score test is that we can do adjustment...
a2 <- qtscore(bt~age+sex,data=srdta,snps=c(1:100),trait.type="binomial")
points(a2[,"Position"],-log10(a2[,"P1df"]),col="green")
```

qvaluebh95

Computes Benjamini-Hochberg (95) q-value

Description

Computes Benjamini-Hochberg (95) q-value

Usage

```
qvaluebh95(p, fdrate=0.1)
```

Arguments

p vector containing p-values

fdrate desired FRD

Value

A list containing

pass Is true if this P-value passed specified FDR

qvalue qvalue

Author(s)

Yurii Aulchenko

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Examples

```
require(GenABEL.data)
data(srdta)
a<-qtscore(qt2,data=srdta)
qv <- qvaluebh95(a[,"P1df"])
plot(a[,"Position"],-log10(qv$qvalue))</pre>
```

r2fast

Estimates r2 between multiple markers

Description

Given a set of SNPs, computes a matrix of r2

Usage

```
r2fast(data, snpsubset, cross.snpsubset, idsubset)
```

Arguments

data object of snp.data-class

snpsubset Index, character or logical vector with subset of SNPs to run analysis on. If

missing, all SNPs from data are used for analysis.

cross.snpsubset

Parameter allowing parallel implementation. Not to be used normally. If sup-

plied together with snpsubset, the r2 for all pairs between snpsubset and cross.snpsubset

computed.

idsubset Index, character or logical vector with subset of IDs to run analysis on. If miss-

ing, all people from data are used for analysis.

Details

The function is based on slightly modified code of Hao et al.

Value

A (Nsnps X Nsnps) matrix giving r2 values between a pairs of SNPs above the diagonal and number of SNP genotype measured for both SNPs below the diagonal

Author(s)

Yurii Aulchenko

References

Hao K, Di X, Cawley S. (2006) LdCompare: rapid computation of single- and multiple-marker r2 and genetic coverage. Bioinformatics, 23: 252-254.

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See Also

rhofast

Examples

```
require(GenABEL.data)
data(ge03d2)
# r2s using r2fast
a <- r2fast(ge03d2,snps=c(1:10))
## Not run:
library(genetics)
# r2s using package genetics
b <- LD(as.genotype(ge03d2[,1:10]))$"R^2"
# see that the r2s are not exactly the same
cor(a[upper.tri(a)],b[upper.tri(b)])
plot(a[upper.tri(a)],b[upper.tri(b)])
## End(Not run)</pre>
```

r2fast.old

Estimates r2 between multiple markers

Description

Given a set of SNPs, computes a matrix of r2

Usage

```
r2fast.old(data, snpsubset, idsubset)
```

Arguments

data object of snp.data-class

snpsubset Index, character or logical vector with subset of SNPs to run analysis on. If

missing, all SNPs from data are used for analysis.

idsubset Index, character or logical vector with subset of IDs to run analysis on. If miss-

ing, all people from data are used for analysis.

Details

The function is based on slightly modified code of Hao et al.

Value

A (Nsnps X Nsnps) matrix giving r2 values between a pairs of SNPs above the diagonal and number of SNP genotype measured for both SNPs below the diagonal

recodeChromosome 107

Author(s)

Yurii Aulchenko

References

Hao K, Di X, Cawley S. (2006) LdCompare: rapid computation of single- and multiple-marker r2 and genetic coverage. Bioinformatics, 23: 252-254.

See Also

rhofast

Examples

```
require(GenABEL.data)
data(ge03d2)
# r2s using r2fast.old
a <- r2fast.old(ge03d2,snps=c(1:10))
## Not run:
library(genetics)
# r2s using package genetics
b <- LD(as.genotype(ge03d2[,1:10]))$"R^2"
# see that the r2s are not exactly the same
cor(a[upper.tri(a)],b[upper.tri(b)])
plot(a[upper.tri(a)],b[upper.tri(b)])
## End(Not run)</pre>
```

recodeChromosome

Change chromosomal coding

Description

Recoding of chromosomes according to the provided 'rules' for from -> to pairs. Most common use is anticipated when importing data from other software using only integers to represent chromosomes. In this situation the list of rules may look like this: list(24="X",25="Y",26="mt").

Usage

```
recodeChromosome(data, rules, quiet = FALSE)
```

Arguments

data	object of class for which 'chromosome' method is defined, e.g. 'gwaa.data', 'snp.data', 'scan.gwaa'
rules	list of pairs 'from=to'; the chromosomes coded in the original data set with 'from' will be recoded with 'to' value
quiet	if summary of recoding should not be printed to the screen

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Value

modified 'data' object

Note

'from' entries should be unique and not overlap with entries in 'to'

Author(s)

Yurii Aulchenko

Examples

```
require(GenABEL.data)
data(ge03d2)
table(chromosome(ge03d2))
# merge chromosome 3 and X, call chromosome 2 as 15
newdat <- recodeChromosome(ge03d2,rules=list("3"="X","2"=15))
table(chromosome(ge03d2),chromosome(newdat))</pre>
```

reconstructNPs

reconstruct nuclear families

Description

Function for reconstruction of nulecar families or extended pedigrees based on results of findRelatives output (estimated meiotic distance matrix). Reconstruction is based on the fact that parent-offspring pairs have meiotic distance of '1', and sibs have a distance '2+2'. If both parents and the offspring are genotyped, expected distance between offspring and both parents is '1', and distance between two parents is >2 (coded as 'NA').

Usage

```
reconstructNPs(relationshipGuessMatrix, sex)
```

Arguments

```
relationshipGuessMatrix
meiotic relationship matrix, as estimated by findRelatives
sex
Sex, coded with 1 for males and 0 for females
```

Value

A matrix containing reconstructed pedigree(s) coded in linkage-like format. If "fid" is zero, this means that a pedigree could not be assigned.

Author(s)

Yurii Aulchenko

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Examples

```
nloci <- 100
q <- runif(nloci,min=0.05,max=0.95)</pre>
# g7---g8
    _|_
  # g9 g10---g11
        __|_
        | /\
#
     g12 g13 g14
#
nids < -8
sex <- c(1,0,0,1,0,0,0,0)
names(sex) <- paste("g",c(7:14),sep="")</pre>
gt <- matrix(ncol=nloci,nrow=nids)</pre>
rownames(gt) <- paste("g",c(7:14),sep="")</pre>
gt["g7",] <- rbinom(nloci,2,q)</pre>
gt["g8",] <- rbinom(nloci,2,q)</pre>
gt["g11",] <- rbinom(nloci,2,q)</pre>
gt["g9",] <- generateOffspring(gt["g7",],gt["g8",],q=q)</pre>
gt["g10",] <- generateOffspring(gt["g7",],gt["g8",],q=q)</pre>
gt["g12",] \leftarrow generateOffspring(gt["g10",],gt["g11",],q=q)
\texttt{gt["g13",]} \leftarrow \texttt{generateOffspring(gt["g10",],gt["g11",],q=q)}
gt["g14",] <- gt["g13",]
aa<-findRelatives(gt,q=q,nmei=c(1:2))</pre>
aa$guess
aaPed <- reconstructNPs(aa$guess,sex)</pre>
aaPed
```

redundant

function to do redundancy check

Description

Checks marker redundancy, understood as comcordance between genotypic distributions (including missing values)

Usage

```
redundant(data, pairs = "bychrom", minconcordance = 2.0)
```

Arguments

data gwaa.data or snp.data object

pairs "bychrom" or "all" to check pairs within chromosome only or genome-wide

minconcordance find "redundant" pairs of markers with concordance >= "minconcordance". If

"minconcordance" is more then 1.0, only pairs of markers which are exactly the

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same (independent of coding), including NA pattern, are considered as redundant. If "minconcordance" is <= 1, the concordance rate is computed as percent of genotypes which are the same, including the genotypes with NA. I.e. if both genotypes are NA, this is counted as a match, if one is NA and other is measured, this is counted as dismatch. Note that option with "minconcordance" <= 1 takes much longer time to run.

Value

A list containing reference SNP as a name and all SNPs which has "the same" genotypic distribution as values:

```
"refSNP1" SNP11, SNP12, ...
"refSNP2" SNP21, SNP22, ...
... etc.
"refSNP1ast" SNP1ast1, SNP1ast2, ...
"all" list of all redundant SNPs, which can be dropped from consideration
```

Author(s)

Yurii Aulchenko

See Also

check.marker

Examples

```
require(GenABEL.data)
data(srdta)
redundant(srdta@gtdata)
redundant(srdta@gtdata[,1:50],minconcordance=0.8)
```

refresh.gwaa.data

Updates an object from old to new GenABEL format

Description

Attempts to update an object of gwaa.data-class from old to new format

Usage

```
refresh.gwaa.data(data,force=FALSE)
```

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Arguments

data An object of gwaa. data-class in pre-1.2-6 (data version 0) format.

force When TRUE, the refreshing is forced, with any data in @strand and @coding

replaced by default data (0/1 coding, u-strand)

Details

Takes old-style gwaa.data object and sets @coding and @strand attributes to SNPs. All coding is set to 1/2 and strand is set to "u" (unknown).

Value

Object of gwaa.data-class in new (GenABEL v > 1.2-6, raw data format version 0.1) format.

Author(s)

Yurii Aulchenko

See Also

load.gwaa.data

rhofast

Estimates rho between multiple markers

Description

Given a set of SNPs, computes a matrix of rho

Usage

```
rhofast(data, snpsubset, idsubset)
```

Arguments

data object of snp.data-class

snpsubset Index, character or logical vector with subset of SNPs to run analysis on. If

missing, all SNPs from data are used for analysis.

idsubset Index, character or logical vector with subset of IDs to run analysis on. If miss-

ing, all people from data are used for analysis.

Details

Rho is the measure of association described by N. Morton and A. Collins (see reference). The function is based on slightly modified code of Hao et al.

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Value

A (Nsnps X Nsnps) matrix giving rho values between a pairs of SNPs above the diagonal and Kij below the diagonal.

Author(s)

Yurii Aulchenko

References

Collins A, Morton NE. (1998) Mapping a disease locus by allelic association. PNAS, 17:1741-1745.

Hao K, Di X, Cawley S. (2006) LdCompare: rapid computation of single- and multiple-marker rho and genetic coverage. Bioinformatics, 23: 252-254.

See Also

r2fast

Examples

```
require(GenABEL.data)
data(ge03d2)
# rhos using rhofast
a <- rhofast(ge03d2,snps=c(1:10))
## Not run:
library(genetics)
# rhos using package genetics
b <- LD(as.genotype(ge03d2[,1:10]))$"R^2"
# see that the rhos are not exactly the same
cor(a[upper.tri(a)],b[upper.tri(b)])
plot(a[upper.tri(a)],b[upper.tri(b)])
## End(Not run)</pre>
```

rntransform

Rank-transformation to normality

Description

Rank-transformation to normality of a variable or residuals from GLM analysis.

Usage

```
rntransform(formula,data,family=gaussian)
```

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Arguments

formula GLM formula for the variable to be transformed, or just the variable

data data.frame or gwaa.data object containing the data

family GLM family

Details

Rank-transformation to normality generates perfectly normal distribution from ANY distribution, unless many/heavy ties are present in variable (or residuals, if formula is used).

When formula is supplied, this procedure first calls ztransform, and then applies rank transformation to residuals.

Value

Vector containing transformed variable, distributed as standard normal.

Author(s)

Yurii Aulchenko

See Also

ztransform

```
# uniformly distributed variable
x <- round(runif(200)*100)</pre>
# get 7 missing values
x[round(runif(7,min=1,max=100))] <- NA
# Z-transform
y0 <- ztransform(x)</pre>
# Rank-transform to normality
y1 <- rntransform(x)</pre>
# test normality of the original and transformed var
shapiro.test(x)
shapiro.test(y0)
shapiro.test(y1)
# plot histogram
par(mfcol=c(3,1))
hist(x)
hist(y0)
hist(y1)
```

114 save.gwaa.data

		.1 - 4 -
save.	gwaa	. gata

function to save gwaa.data object

Description

Saves GenABEL data in internal format

Usage

```
save.gwaa.data(data, phenofile = "pheno.dat", genofile = "geno.raw",
human = FALSE)
```

Arguments

data gwaa.data object

phenofile name of file where the phenotypes will be saved to genofile name of file where the genotypes will be saved to

human if human=TRUE, saves in human-readable format (to be converted to internal

format later)

Details

When running with human=TRUE, a lot of memory (and time to complete the operation) is required. Probably, this option would not work because of memory limitations in a GWA scan iwth more then few hundreds of people. This is possible to fix; drop me a message if you need that.

Value

No value returned

Author(s)

Yurii Aulchenko

See Also

load.gwaa.data

scan.glm 115

	-	
scan	.glm	

Scan GWA data using glm

Description

Scan GWA data using glm

Usage

```
scan.glm(formula, family = gaussian(), data, snpsubset, idsubset,
bcast = 50)
```

Arguments

formula character string containing formula to be used in glm. You should put CRSNP

argument in the formula, to arrange how the SNP from the list would be treated.

This allows to put in an interaction term.

family family to be passed to glm

snpsubset Index, character or logical vector with subset of SNPs to run analysis on. If

missing, all SNPs from data are used for analysis.

idsubset Index, character or logical vector with subset of IDs to run analysis on. If miss-

ing, all people from data/cc are used for analysis.

data object of class "gwaa.data"

bcast show progress every bcast SNPs

Value

```
Object of class scan.gwaa-class
```

Author(s)

Yurii Aulchenko

See Also

```
ccfast, qtscore, scan.gwaa-class
```

```
require(GenABEL.data)
data(srdta)
a <- scan.glm("bt~sex+age+CRSNP",family=binomial(),data=srdta,snps=(1:10),bcast=2)
#plot(a)

osnp <- "rs4934"
maposnp <- srdta@gtdata@map[osnp]
maposnp</pre>
```

scan.glm.2D

```
reg <- snp.names(srdta,begin=maposnp-100000,end=maposnp+100000,chrom="1")
## Not run:
a <- scan.glm("qt3~sex+age+CRSNP",data=srdta,snps=reg)
plot(a)
plot(a,df=1)
add.plot(a,df=2)
## End(Not run)

# interaction with sex
## Not run:
a <- scan.glm("qt3~age+sex*CRSNP",data=srdta,snps=reg)
plot(a,df=1)
add.plot(a,df=2)
# you can do interaction with a selected polymorphisms in the same way
## End(Not run)</pre>
```

scan.glm.2D

Scans regional data allowing for gene-gene interaction using glm

Description

Scans regional data allowing for gene-gene interaction using glm

Usage

```
scan.glm.2D(formula, family = gaussian(), data, snpsubset, idsubset,
bcast = 50)
```

Arguments

formula	character string containing formula to be used in glm. You should put CRSNP argument in the formula, to arrange how the SNP from the list would be treated. This allows to put in an interaction term.
family	family to be passed to glm
snpsubset	Index, character or logical vector with subset of SNPs to run analysis on. If missing, all SNPs from data are used for analysis.
idsubset	Index, character or logical vector with subset of IDs to run analysis on. If missing, all people from data/cc are used for analysis.
data	object of class "gwaa.data"
bcast	show progress every bcast SNPs

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Details

For each pair of SNPs, say snp1 and snp2, scan.glm.2D estimates 5 models. Let us denote snp1 when it is coded as allele dose (0,1, 2) and thus results in additive model as snp1dose and when it is coded as 'factor' (genotypic model) as snp1factor

```
m00: y ~ mu [1 regression coefficient to estimate]
m10: y ~ mu + snp1dose + snp2dose [3 coefficients]
m11: y ~ mu + snp1dose + snp2dose + snp1dose * snp2dose [4 coefficients]
m20: y ~ mu + snp1factor + snp2factor [5 coefficients]
m21: y ~ mu + snp1factor + snp2factor + snp1factor * snp2factor [9 coefficients]
```

In the output, "P1df" refers to the test of m00 vs m10 (this is actually 2 df test); "P2df" refers to the test of m00 vs m20 (4 df test); "Pint1df" refers to the test of m10 vs m11 (1 df test); "Pint2df" refers to the test of m20 vs m21 (4 df test). The output is in matrix format as these P-values are generated for each pair of SNPs in turn.

Value

```
Object of class scan.gwaa.2D-class
```

Author(s)

Yurii Aulchenko

See Also

```
scan.gwaa.2D-class, scan.haplo.2D
```

Examples

```
## Not run:
require(GenABEL.data)
data(srdta)
a <- scan.glm.2D("bt~sex+age+CRSNP",family=binomial(),data=srdta,snps=(1:10),bcast=2)
plot(a)
## End(Not run)</pre>
```

scan.gwaa-class

Class "scan.gwaa"

Description

This class contains results of GWA analysis. This is an list object, generated by scan.glm, scan.haplo, ccfast, qtscore, emp.ccfast, or emp.qtscore.

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Names

snpnames list of names of SNPs tested

P1df corresponding list of P-values of 1-d.f. (additive or allelic) test for association bestween SNP and trait

P2df corresponding list of P-values of 2-d.f. (genotypic) test for association bestween SNP and trait

Pc1df P-values from the 1-d.f. test for association bestween SNP and trait; the statistics is corrected for possible inflation

effB Effect of the B allele in allelic test (OR for ccfast, difference from the mean for qtscore and beta from the scan.glm)

effAB Effect of the AB genotype in genotypic test

effBB Effect of the BB genotype in genotypic test

map list of map positions of the SNPs

chromosome list of chromosomes the SNPs belong to

idnames list of people used in analysis

lambda list with elements "estimate" (inflation factor estimate, as computed using lower 90 percents of the distribution) and "se" (standard error of the estimate)

formula which formula/function call was used to comput P-values

family family of the link function / nature of the test

Methods

Author(s)

Yurii Aulchenko

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See Also

ccfast, qtscore, scan.glm, scan.haplo, emp.ccfast, emp.qtscore, estlambda, plot.scan.gwaa

Examples

```
require(GenABEL.data)
data(srdta)
sc <- qtscore(qt3,data=srdta,snps=c(1:10))
class(sc)
sc[,"P1df"]
sc[,"P2df"]
sc
plot(sc)</pre>
```

scan.gwaa.2D-class

Class "scan.gwaa.2D"

Description

This class contains results of 2D analysis. This is an list object, generated by scan.glm.2D or scan.haplo.2D.

Names

snpnames list of names of SNPs tested

P1df corresponding list of P-values of allelic test for association bestween SNP and trait.

Pint1df corresponding list of P-values of significance of the interactions between SNPs, for the allelic model

P2df corresponding list of P-values of genotypic test for association bestween SNP and trait For link{scan.haplo} and link{scan.haplo.2D} this is equal to P1df and has nothing to do with the actual degrees fo freedom of the test

Pint1df corresponding list of P-values of significance of the interactions between SNPs for the genotypic test

medChi1df Median Chi-square for allelic test

medChi2df Median Chi-square on genotypic test

map list of map positions of the SNPs

chromosome list of chromosomes the SNPs belong to

formula which formula/function call was used to comput P-values

family family of the link function / nature if the test

idnames list of people used in analysis

Methods

plot signature(object = "scan.gwaa.2D"): Plots summary of 2D scan, using lsit element P1df

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Author(s)

Yurii Aulchenko

See Also

```
scan.gwaa.2D-class, scan.glm.2D, scan.haplo.2D, plot.scan.gwaa.2D
```

Examples

```
require(GenABEL.data)
data(srdta)
sc <- scan.glm.2D("qt3~CRSNP",data=srdta,snps=c(1:10))
class(sc)
sc$P1df
sc$P2df
sc
plot(sc)</pre>
```

scan.haplo

scan.haplo

Description

Runs haplo.score.slide from the package haplo.stats and represents output as scan.gwaa-class data object

Usage

```
scan.haplo(formula, data, snpsubset, idsubset, n.slide = 2, bcast = 10,
simulate=FALSE, trait.type, ...)
```

Arguments

Formula to be used in analysis. It should be a character string following standard notation. On the left-had side, there should be outcome. On the right-hand side,

covariates are liste, with "+" separating the covariates (additive action). The leftand right-hand sides are separated by "~". You should put CRSNP argument in the formula. For example "qt3~CRSNP" would analyse association between SNPs and trait "qt3", without any adjustment. To adjust for age and sex, use "qt3~age+sex+CRSNP". Currently, only additive effects ("+") are allowed.

data object of calss gwaa.data-class

snpsubset Index, character or logical vector with subset of SNPs to run analysis on. If

missing, all SNPs from data are used for analysis.

idsubset Index, character or logical vector with subset of IDs to run analysis on. If miss-

ing, all people from data/cc are used for analysis.

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n.slide	Default = 2. Number of loci in each contiguous subset. The first subset is the ordered loci numbered 1 to n.slide, the second subset is 2 through n.slide+1 and so on. If the total number of loci in geno is n.loci, then there are n.loci - n.slide + 1 total subsets.
bcast	show progress every bcast SNPs
simulate	if simulated P-values should be generated
trait.type	Character string defining type of trait, with values of "gaussian", "binomial", "poisson", "ordinal" (see help for haplo.score.slide for details). If not specified, the routine picks up "gaussian" or "binomial" (two levels of trait).
	other arguments to be passed to haplo.score.slide

Details

List element P2df is set equal to P1df, as only allelic results are returned. This has nothing to do with degrees of freedom.

Value

```
Object of class scan.gwaa-class
```

Author(s)

Yurii Aulchenko

References

For haplo.stats (scan.haplo, scan.haplo.2D), please cite:

Schaid DJ, Rowland CM, Tines DE, Jacobson RM, Poland GA. (2002) Score tests for association between traits and haplotypes when linkage phase is ambiguous. Am J Hum Genet, 70: 425-434.

See Also

```
scan.gwaa-class, haplo.score.slide
```

```
require(GenABEL.data)
data(srdta)
a <- ccfast("bt",srdta,snps=(717:733),ids=(srdta@phdata$age<40))
plot(a)
if (require(haplo.stats)) {
b <- scan.haplo("bt~sex+CRSNP",srdta,snps=(717:733),
ids=(srdta@phdata$age<40))
c <- scan.haplo("bt~sex+CRSNP",srdta,snps=(717:733),
ids=(srdta@phdata$age<40),n.slide=3)
add.plot(b,col="red",type="l")
add.plot(c,col="darkgreen",type="l")
}</pre>
```

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scan.haplo.2D

runs haplo.score.slide with all pairs of markers in a region

Description

Runs haplo.score.slide from the package haplo.stats on all pairs of markers in a region and presents output as scan.gwaa.2D-class object

Usage

```
scan.haplo.2D(formula, data, snpsubset, idsubset, bcast = 10,
simulate=FALSE, trait.type, ...)
```

Arguments

formula	Formula to be used in analysis. It should be a character string following standard notation. On the left-had side, there should be outcome. On the right-hand side, covariates are liste, with "+" separating the covariates (additive action). The left-and right-hand sides are separated by "~". You should put CRSNP argument in the formula. For example "qt3~CRSNP" would analyse asociation between SNPs and trait "qt3", without any adjustment. To adjust for age and sex, use "qt3~age+sex+CRSNP". Currently, only additive effects ("+") are allowed.
data	object of calss gwaa.data-class
snpsubset	Index, character or logical vector with subset of SNPs to run analysis on. If missing, all SNPs from data are used for analysis.
idsubset	Index, character or logical vector with subset of IDs to run analysis on. If missing, all people from data/cc are used for analysis.
bcast	show progress every bcast percents of progress
simulate	if simulated P-values should be generated
trait.type	Character string defining type of trait, with values of "gaussian", "binomial", "poisson", "ordinal" (see help for haplo.score.slide for details). If not specified, the routine picks up "gaussian" or "binomial" (two levels of trait).
	other arguments to be passed to haplo.score.slide

Details

List element P2df is set equal to P1df, as only allelic results are returned. This has nothing to do with actual degrees of freedom of the test.

Value

```
Object of class scan.gwaa.2D-class
```

Author(s)

Yurii Aulchenko

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References

For haplo.stats (scan.haplo, scan.haplo.2D), please cite:

Schaid DJ, Rowland CM, Tines DE, Jacobson RM, Poland GA. (2002) Score tests for association between traits and haplotypes when linkage phase is ambiguous. Am J Hum Genet, 70: 425-434.

See Also

```
scan.gwaa.2D-class, scan.haplo, scan.glm.2D, haplo.score.slide
```

Examples

```
if (require(haplo.stats)) {
## Not run:
require(GenABEL.data)
data(srdta)
c <- scan.haplo.2D("bt~sex+age+CRSNP",data=srdta,snps=(717:733),ids=(srdta@phdata$age<40))
plot(c)
## End(Not run)
}</pre>
```

show.ncbi

Shows the region on NCBI map

Description

This function calls web browser and direct it to NCBI MapViewer, to show the region of interest.

Usage

```
show.ncbi(region)
```

Arguments

region

a vector containing regional landmarks

Details

The elements of input vector could be SNP rs-names

Author(s)

Yurii Aulchenko

124 snp.coding-class

Examples

```
## Not run:
show.ncbi(c("rs7926624","rs11564708"))
## End(Not run)
```

snp.coding-class

Class "snp.coding"

Description

This class contains the actual nucleotide codes for the typed SNPs

Slots

.Data: nucleotide coding data

Methods

```
[ signature(x = "snp.coding", i = "ANY", j = "missing", drop = "missing"): subset operations. x[i] will show coding for SNPs selected in i.
```

coerce signature(from = "snp.coding", to = "character"): converts SNP coding from
 internal (raw) to human-readable character.

show signature(object = "snp.coding"): shows the object. Take care that this is internal
representation

Author(s)

Yurii Aulchenko

See Also

```
snp.strand-class, gwaa.data-class, snp.data-class
```

```
require(GenABEL.data)
data(srdta)
srdta@gtdata@coding[1:10]
as.character(srdta@gtdata@coding[1:10])
```

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snp.data

creates an snp.data object

Description

Creates object of class snp.data-class

Usage

```
snp.data(nids, rawdata, idnames = as.character(c(1:nids)),
snpnames = as.character(c(1:(length(rawdata)/ceiling(nids/4)))),
chromosome = as.factor(rep(1,(length(rawdata)/ceiling(nids/4)))),
map = as.double(seq(1,(length(rawdata)/ceiling(nids/4)))),
coding=as.raw(rep(1,length(rawdata)/ceiling(nids/4))),
strand=as.raw(rep(0,length(rawdata)/ceiling(nids/4))),
male = rep(0, nids))
```

Arguments

nids number of people

idnames list of IDs

male male indicator for IDs

snpnames list of SNP names

chromosome list of chromosomes SNPs belong to coding list of nucleotide coding for the SNPs

strand strands of the SNPs
map map position of SNPs

rawdata genotypes presented in raw data format

Value

Object of class snp.data-class

Author(s)

Yurii Aulchenko

See Also

```
snp.data-class
```

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snp.data-class

Class "snp.data"

Description

This class contains objects holding large arrays of single nucleotide polymorphism (SNP) genotypes

Slots

```
nbytes: number of bytes used to store data on a SNP
nids: number of people
male: male code
idnames: ID names
nsnps: number of SNPs
snpnames: list of SNP names
chromosome: list chromosomes corresponding to SNPs
coding: list of nucleotide coding for the SNPs
strand: strands of the SNPs
map: list SNPs' positions
gtps: snp.mx-class object used to store genotypes
```

Methods

```
[ signature(x = "snp.data", i = "ANY", j = "ANY", drop = "ANY"): subset operations.
     x[i,j] will select people listed in i and SNPs listed in j.
coerce signature(from = "snp.data", to = "numeric"): map to codes 0, 1, 2, or NA
coerce signature(from = "snp.data", to = "character"): map to actual nucleotide codes,
    e.g. "A/A", "A/G", "G/G", ""
coerce signature(from = "snp.data", to = "genotype"): map to data frame with genotype-
    class data, for later use with package genetics
coerce signature(from = "snp.data", to = "hsgeno"): map to data frame with allelic data
     frame, for later use with package haplo.stats
show signature(object = "snp.data"): shows the object. Take care that the objects are usually
     very large!
summary signature(object = "snp.data"): calculate allele frequencies, genotype frequen-
    cies, and chi-square tests for Hardy-Weinberg equilibrium. Results are returned as a dataframe
annotation signature(object = "gwaa.data"), signature(object = "snp.data"): extracts
     annotation
idnames signature(object = "gwaa.data"), signature(object = "snp.data"): extracts id
    names
```

snp.data-class 127

```
snpnames signature(object = "gwaa.data"), signature(object = "snp.data"): extracts
    snp names
nids signature(object = "gwaa.data"), signature(object = "snp.data"): extracts num-
    ber of ids
nsnps signature(object = "gwaa.data"), signature(object = "snp.data"): extracts num-
    ber of snps
map signature(object = "gwaa.data"), signature(object = "snp.data"): extracts map
chromosome signature(object = "gwaa.data"), signature(object = "snp.data"): ex-
    tracts chromosome
strand signature(object = "gwaa.data"), signature(object = "snp.data"): extracts
    strand
strand<- signature(object = "gwaa.data"), signature(object = "snp.data"): assign</pre>
coding signature(object = "gwaa.data"), signature(object = "snp.data"): extracts
    coding
coding<- signature(object = "gwaa.data"), signature(object = "snp.data"): assign</pre>
    coding
refallele signature(object = "gwaa.data"), signature(object = "snp.data"): extracts
    reference allele
effallele signature(object = "gwaa.data"), signature(object = "snp.data"): extracts
    effective allele
male signature(object = "gwaa.data"), signature(object = "snp.data"): extracts male
    indicator
```

Author(s)

Yurii Aulchenko

See Also

```
gwaa.data-class, snp.data, snp.mx-class
```

```
require(GenABEL.data)
data(srdta)
class(srdta)
x <- srdta@gtdata
class(x)
nids(x)
nsnps(x)
idnames(x)[1:12]
male(x)[1:12]
male(x)[c("p1","p2","p3","p4")]
snpnames(x)[1:4]
chromosome(x)[1:4]
map(x)[1:4]</pre>
```

128 snp.mx-class

```
n4 <- c("rs18","rs655")
n4
map(x)[n4]
n4 <- c("rs18","rs65")
n4
map(x)[n4]
chromosome(x)[n4]
x[1:12,1:4]
summary(x[,1:10])
as.numeric(x[1:12,1:4])
as.numeric(x[c("p1","p3","p4"),c("rs18","rs65")])
as.character(x[c("p1","p3","p4"),c("rs18","rs65")])
as.genotype(x[c("p1","p3","p4"),c("rs18","rs65")])
as.hsgeno(x[c("p1","p3","p4"),c("rs18","rs65")])</pre>
```

snp.mx-class

Class "snp.mx"

Description

This low-level class contains objects holding large arrays of single nucleotide polymorphism (SNP) genotypes

Slots

.Data: object used to store genotypes

Methods

```
[ signature(x = "snp.mx", i = "ANY", j = "ANY", drop = "ANY"): subset operations.
    x[i,j] will select people listed in i and SNPs listed in j.

coerce signature(from = "raw", to = "snp.mx"): makes an snp.mx object out of raw data
show signature(object = "snp.mx"): shows the object. Take care that (a) this is internal
    representation and (b) the objects are usually very large!
```

Note

User is not supposed to work with this class. Use snp.data-class.

Author(s)

Yurii Aulchenko

See Also

```
gwaa.data-class, snp.data-class
```

snp.names 129

snp.names	extracts names of SNPs in a region	

Description

Based on boundary conditions specified and (or) chromosome selects SNP names in the region

Usage

```
snp.names(data, begin, end, chromosome)
```

Arguments

data object of class gwaa.data-class, snp.data-class, scan.gwaa-class or check.marker-class

begin Start position (or name of the first SNP)
end End-position or name of last SNP

chromosome Chromosome code

Details

Any of the arguments, except the data can be missing

Value

A vector of names of SNPs located in the region

Author(s)

Yurii Aulchenko

See Also

```
snp.data-class
```

```
require(GenABEL.data)
data(srdta)
snp.names(srdta, begin = 50000, end = 100000)
snp.names(srdta, begin = 50000, end = 100000, chromosome = "1")
# does not make sense with these data:
snp.names(srdta, begin = 50000, end = 100000, chromosome = "X")
# again makes sense:
snp.names(srdta, end = 100000)
snp.names(srdta, begin = 2200000)
```

snp.strand-class

```
# show summary for SNPs in region between 50,000 and 100,000
a <- snp.names(srdta, begin = 50000, end = 100000)
summary(srdta@gtdata[,a])</pre>
```

snp.strand-class

Class "snp.strand"

Description

This class contains the strands of the typed SNPs

Slots

.Data: nucleotide strand data

Methods

```
[ signature(x = "snp.strand", i = "ANY", j = "missing", drop = "missing"): subset operations. x[i] will show strand for SNPs selected in i.
```

coerce signature(from = "snp.strand", to = "character"): converts SNP strand from internal (raw) to human-readable character.

show signature(object = "snp.strand"): shows the object. Take care that this is internal
representation

Author(s)

Yurii Aulchenko

See Also

```
snp.coding-class, gwaa.data-class, snp.data-class
```

```
require(GenABEL.data)
data(srdta)
srdta@gtdata@strand[1:10]
as.character(srdta@gtdata@strand[1:10])
```

snp.subset 131

snp.subset

function to subset objects of class scan.gwaa and check.marker

Description

Computing objects of class scan.gwaa may take long, especially when haplotypic analysis is performed. Therefore this function helps substracting results on some region (indicated by list of SNPs)

Usage

```
snp.subset(data, snpsubset)
```

Arguments

data object of class scan.gwaa-class or check.marker-class

snpsubset character vector of snps to select

Value

Object of class scan.gwaa-class or check.marker-class

Author(s)

Yurii Aulchenko

See Also

```
scan.gwaa-class, check.marker-class
```

```
require(GenABEL.data)
data(srdta)
# processing check.marker object
#mc <- check.marker(data=srdta@gtdata[,1:100],redundant="all",maf=0.01,
# minconcordance=0.9,fdr=.1,ibs.mrk=0)
mc <- check.marker(data=srdta@gtdata[,1:100],maf=0.01,fdr=.1,ibs.mrk=0)
summary(mc)
#plot(mc)
mc1 <- snp.subset(mc,snps=srdta@gtdata@snpnames[20:50])
summary(mc1)
#plot(mc1)
# processing scan.gwaa object
a <- qtscore(qt3~sex+age,data=srdta)
plot(a)
a1 <- snp.subset(a,snps=srdta@gtdata@snpnames[10:20])
plot(a1)</pre>
```

132 sortmap.internal

snps.cell-class

Class "snps.cell"

Description

This is a lowest-level class based on which snp.mx-class is build

Note

User is not supposed to work with this class. Use snp.data-class.

Author(s)

Yurii Aulchenko

See Also

```
snp.mx-class, gwaa.data-class, snp.data-class
```

sortmap.internal

Internal function for map-sorting

Description

Internal function for map-sorting, not supposed to be used directly by user (is open for regression testing reasons)

Usage

```
sortmap.internal(chrom, map, delta = 1)
```

Arguments

chrom vector of markers' chromosomes
map vector of marlers' map ositions

delta step to do between chroms when building cumulative map

Value

list, withe elements 'ix' ('sorted' order), etc.

Author(s)

Yurii Aulchenko

sset 133

sset

Internal use function for class snp.mx-class

Description

Interface to C function sset subsetting genotypes from snp.mx-class

Usage

```
sset(data, nsnps, nids, list)
```

Arguments

data genotypic data in internal format

nsnps no. snps nids no. people

list something internal...

Details

Rather simple function which I wrote before discovering R's setdiff, etc. functions.

Value

Sub-set from snp.mx-class object

Author(s)

Yurii Aulchenko

See Also

```
snp.mx-class
```

summary.check.marker

Summary of check.marker object

Description

Provides cross-tabulation summarising number of marker which did not pass this or that criteria

Usage

```
## S3 method for class 'check.marker'
summary(object, ...)
```

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Arguments

```
object of class check.marker-class
... additional arguments (not used)
```

Value

A list containing 2 tables: per-marker and per-person inconsistencies

Author(s)

Yurii Aulchenko

See Also

```
check.marker, check.marker-class
```

Examples

```
require(GenABEL.data)
data(srdta)
mc <- check.marker(srdta,ids=c(1:500))
summary(mc)</pre>
```

summary.gwaa.data

function to summarise GWAA data

Description

Summary of phenotypic and genotypic parts of GWAA data

Usage

```
## S3 method for class 'gwaa.data'
summary(object, ...)
```

Arguments

```
object of class gwaa.data-class
... additional arguments (not used)
```

Value

Returns list with two elements:

pheno Summary for phenotypic part of gwaa.data object geno Summary for genotypic part of gwaa.data object

summary.scan.gwaa 135

Author(s)

Yurii Aulchenko

See Also

```
summary.snp.data
```

Examples

```
require(GenABEL.data)
data(srdta)
# be prepared : long output!
summary(srdta)
```

summary.scan.gwaa

Shortcut to 'descriptives...

Description

Shortcut to 'descriptives.scan'

Usage

```
## S3 method for class 'scan.gwaa'
summary(object, ...)
```

Arguments

```
object of class 'scan.gwaa' as generated by qtscore, mlreg, etc.
... arguments passed to descriptives.scan
```

Details

Shortcut to 'descriptives.scan'

Author(s)

Yurii Aulchenko

```
require(GenABEL.data)
data(srdta)
x <- qtscore(qt2,srdta)
summary(x)</pre>
```

136 summary.snp.data

summary.snp.data

function to summary GWAA data

Description

Provides summary of an object of class snp.data-class. Chromosome, map position, allele coding, number of observed genotypes, allelic frequency, genotypic distribution, P-value of the exact test for HWE, Fmax (estimate of deviation from HWE, allowing meta-analysis) and LRT P-value for HWE test are listed

Usage

```
## S3 method for class 'snp.data'
summary(object, ...)
```

Arguments

object snp.data object

... additional arguments (not used)

Value

Data frame summary for snp.data object

Note

The P-values reported for X-chromosome are based on analysis of female data, but other statistics (frequencies, calls, ...) are based on all data. Statistics for Y-chromosome are based on male-only. P-HWE is not defined for mt- and Y- markers (set to 1.0).

Author(s)

Yurii Aulchenko

References

Wigginton, JE, Cutler, DJ, and Abecasis, GR (2005) A Note on Exact Tests of Hardy-Weinberg Equilibrium. American Journal of Human Genetics. 76: 887-93.

See Also

```
summary.gwaa.data, snp.data-class
```

```
require(GenABEL.data)
data(srdta)
summary(gtdata(srdta[,1:20]))
```

VIFGC 137

VIFGC	Genomic control for varioi	us model of inheritance using VIF
-------	----------------------------	-----------------------------------

Description

This function estimates corrected statistic using genomic control for different models (recessive, dominant, additive etc.), using VIF. VIF coefficients are estimated by optimizing different error functions: regress, median and ks.test.

Usage

```
VIFGC(data, p, x, method = "regress", n,
  index.filter = NULL, proportion = 1, clust = 0,
  vart0 = 0, tmp = 0, CA = FALSE, p.table = 0,
  plot = TRUE, lmax = NULL, color = "red", F = NULL,
  K = NULL, type_of_plot = "plot", ladd = NULL)
```

Arguments

data	Input vector of Chi square statistic
method	Function of error to be optimized. Can be "regress", "median" or "ks.test"
р	Input vector of allele frequencies
X	Model of inheritance (0 for recessive,0.5 for additive, 1 for dominant, also it could be arbitrary)
index.filter	Indexes for variables that will be use for analysis in data vector
n	The size of the sample
proportion	The proportion of lowest P (Chi2) to be used when estimating the inflation factor Lambda for "regress" method only
plot	If TRUE, plot of lambda will be produced
type_of_plot	For developers only
lmax	The threshold for lambda for plotting (optional)
color	The color of the plot
F	The estimation of F (optional)
K	The estimation of K (optional)
ladd	The estimation of lambda for additive model (optional)
clust	For developers only
vart0	For developers only
tmp	For developers only
CA	For developers only
p.table	For developers only

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Value

A list with elements

Zx output vector corrected Chi square statistic

vv output vector of VIF

exeps output vector of exepsons (NA)

calrate output vector of calrate

F F K

Author(s)

Yakov Tsepilov

Examples

```
require(GenABEL.data)
data(ge03d2)
# truncate the data to make the example faster
ge03d2 <- ge03d2[seq(from=1,to=nids(ge03d2),by=2),seq(from=1,to=nsnps(ge03d2),by=3)]
qts <- mlreg(dm2~sex,data=ge03d2,gtmode = "dominant")
chi2.1df <- results(qts)$chi2.1df
s <- summary(ge03d2)
freq <- s$Q.2
result <- VIFGC(p=freq,x=1,method = "median",CA=FALSE,data=chi2.1df,n=nids(ge03d2))</pre>
```

VIFGC_ovdom

Genomic control for over-dominant model of inheritance using VIF

Description

This function estimates the corrected statistic using genomic control for the over-dominant model, using VIF. VIF coefficients are estimated by optimizing different error functions: regress, median and ks.test.

Usage

```
VIFGC_ovdom(data, p, method = "regress", n,
  index.filter = NULL, proportion = 1, clust = 0,
  vart0 = 0, tmp = 0, plot = TRUE, lmax = NULL,
  color = "red")
```

VIFGC_ovdom 139

Arguments

data Input vector of Chi square statistic

method Function of error to be optimized. Can be "regress", "median" or "ks.test"

p Input vector of allele frequencies

index.filter Indexes for variables that will be use for analysis in data vector

n size of the sample

proportion The proportion of lowest P (Chi2) to be used when estimating the inflation factor

Lambda for "regress" method only

plot If TRUE, plot of lambda will be produced

1max The threshold for lambda for plotting (optional)

color The color of the plot
clust For developers only
vart0 For developers only
tmp For developers only

Value

A list with elements

Zx output vector corrected Chi square statistic

vv output vector of VIF

exeps output vector of exepsons (NA)

calrate output vector of calrate

F F K

Author(s)

Yakov Tsepilov

```
require(GenABEL.data)
data(ge03d2)
# truncate the data to make the example faster
ge03d2 <- ge03d2[seq(from=1,to=nids(ge03d2),by=2),seq(from=1,to=nsnps(ge03d2),by=3)]
qts <- mlreg(phdata(ge03d2)$dm2~1,data=ge03d2,gtmode = "overdominant")
chi2.1df <- results(qts)$chi2.1df
s <- summary(ge03d2)
freq <- s$Q.2
result <- VIFGC_ovdom(p=freq,method = "median",data=chi2.1df,n=nids(ge03d2))</pre>
```

X fix

Xfix

function to set impossible genotypes as missing

Description

Sets impossible genotypes (e.g. heterozygous male X-linked genotypes) to missing

Usage

```
Xfix(data)
```

Arguments

data

Object of gwaa.data-class

Details

Sets to missing genotypes in the following situations: (1) heterozygous male X-genotypes (2) heterozygous Y- and mtDNA genotypes (3) any Y-genotypes in females. Should only be used after check.marker, which identifies systematic sex errors.

Value

The same object of gwaa.data-class, with fixed genotypes

Author(s)

Yurii Aulchenko

See Also

check.marker

```
require(GenABEL.data)
data(ge03d2c)
# truncate the data to make the example faster
ge03d2c <- ge03d2c[seq(from=1,to=nids(ge03d2c),by=2),seq(from=1,to=nsnps(ge03d2c),by=2)]
# many errors
mc0 <- check.marker(ge03d2c)
# take only people and markers passing QC
fixed0 <- ge03d2c[mc0$idok,mc0$snpok]
# major errors fixed, still few males are heterozygous for X-chromsome markers
mc1 <- check.marker(fixed0)
# fix minor X-chromosome problems
fixed1 <- Xfix(fixed0)
# no errors
mc2 <- check.marker(fixed1)
summary(mc2)</pre>
```

ztransform 141

Description

Transformation of a variable or residuals from GLM analysis to standard Normal.

Usage

```
ztransform(formula,data,family=gaussian)
```

Arguments

formula GLM formula for the variable to be transformed, or just the variable

data data.frame or gwaa.data object containing the data

family GLM family

Details

Transformation to normality generates a variable which has mean zero and variance of one. If formula used, residuals from regression model are scaled to standard Normal.

Value

Vector containing transformed variable, distributed as standard normal.

Author(s)

Yurii Aulchenko

See Also

ztransform

```
# uniformly distributed variable
x <- round(runif(200)*100)
# get 7 missing values
x[round(runif(7,min=1,max=100))] <- NA
# Z-transform
y0 <- ztransform(x)
# Rank-transform to normality
y1 <- rntransform(x)
# test normality of the original and transformed var
shapiro.test(x)
shapiro.test(y0)
shapiro.test(y1)</pre>
```

142 ztransform

```
# plot histogram
par(mfcol=c(3,1))
hist(x)
hist(y0)
hist(y1)
# tests with genetic data
require(GenABEL.data)
data(srdta)
Zqt1 <- ztransform(qt1,srdta)
Zqt1sexA <- ztransform(qt1~sex,srdta)</pre>
```

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