

# snpStats

March 24, 2012

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Fst

*Calculate fixation indices*

---

## Description

This function calculates the fixation index  $F_{st}$  for each SNP, together with its weight in the overall estimate (as used by the International HapMap Consortium).

## Usage

```
Fst(snp, group)
```

## Arguments

<code>snp</code>	an object of class <code>SnpMatrix</code> or <code>XSnpMatrix</code> containing the SNP data
<code>group</code>	a factor (or object that can be coerced into a factor), of length equal to the number of rows of <code>snp</code> , giving the grouping or rows for which the $F_{st}$ is to be calculated

## Value

A list:

**Fst**  $F_{st}$  values for each SNP

**weight** The weights for combining these into a single index

## Note

Uncertain genotypes are treated as missing

## Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

**Examples**

```
## Analysis of some HapMap data

data(for.exercise)
f <- Fst(snps.10, subject.support$stratum)
weighted.mean(f$Fst, f$weight)
```

---

GlmEstimates-class *Class "GlmEstimates"*

---

**Description**

A simple class to hold output from `snp.lhs.estimate`s and `snp.rhs.estimate`s. Its main purpose is to provide a `show` method

**Objects from the Class**

Objects from this class are simple lists. Each element of the list is a list giving the results of a generalized linear model fit, with elements:

**Y.var** Name of the Y variable

**beta** The vector of parameter estimates (with their names)

**Var.beta** The upper triangle of the variance-covariance matrix of estimates, stored as a simple vector

**N** The number of "units" used in the model fit

**Extends**

Class "`list`", from data part. Class "`vector`", by class "`list`", distance 2.

**Methods**

[ signature(x = "GlmEstimates", i = "ANY", j = "missing", drop = "missing"): Subset

**coerce** signature(from = "GlmEstimates", to = "GlmTests"): Calculate Wald tests

**show** signature(object = "GlmEstimates"): Display

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**See Also**

[snp.lhs.estimate](#), [snp.rhs.estimate](#)

**Examples**

```
showClass("GlmEstimates")
```

---

GlmTests-class      *Classes "GlmTests" and "GlmTestsScore"*

---

### Description

Classes of objects created by `snp.lhs.tests` and `snp.rhs.tests`. The class "GlmTestsScore" extends the class "GlmTests" and is invoked by setting the argument `score=TRUE` when calling testing functions in order to save the scores and their variances (and covariances)

### Objects from the Class

Objects of class "GlmTests" have four slots:

**snp.names** When only single SNPs are tested, a character vector of SNP names. Otherwise a list of such vectors (one for each test)

**var.names** A character vector containing names of variables tested against SNPs

**chisq** A numerical vector of chi-squared test values

**df** An integer vector of degrees of freedom for the tests

**N** A integer vector of the number of samples contributing to each test

The "GlmTestsScore" class extends this, adding a slot `score` containing a list with elements which are themselves lists with two elements:

**U** The vector (or matrix) of efficient scores

**V** The upper triangle of the variance-covariance matrix of U, stored as a vector

### Methods

[ ] signature(x = "GlmTests", i = "ANY", j = "missing", drop = "missing"): Subsetting operator

**coerce** signature(from = "GlmTests", to = "data.frame"): Simplify object

**chi.squared** signature(x = "GlmTests", df = "missing"): Extract chi-squared test values

**deg.freedom** signature(x = "GlmTests"): Extract degrees of freedom for tests

**names** signature(x="GlmTests"): Extract (or generate) a name for each test

**p.value** signature(x = "GlmTests", df = "missing"): Extract *p*-values

**sample.size** signature(object = "GlmTests"): Extract sample sizes for tests

**show** signature(object = "GlmTests"): Show method

**summary** signature(object = "GlmTests"): Summary method

[ ] signature(x = "GlmTestsScore", i = "ANY", j = "missing", drop = "missing"): Subsetting operator

**effect.sign** signature(x = "GlmTestsScore", simplify = "logical"): Extract signs of associations. If `simplify` is TRUE then a simple vector is returned if all tests are on 1df

**pool2** signature(x = "GlmTestsScore", y = "GlmTestsScore", score = "logical"): Combine results from two sets of tests

**switch.alleles** signature(x = "GlmTestsScore", snps = "character"): Emulate, in the score vector and its (co)variances, the effect of switching of the alleles of specified SNPs

**Note**

Most of the methods for this class are shared with the [SingleSnpTests](#) and [SingleSnpTestsScore](#) classes

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**See Also**

[snp.lhs.tests](#), [snp.rhs.tests](#), [SingleSnpTests](#), [SingleSnpTestsScore](#)

**Examples**

```
showClass("GlmTests")
```

---

```
ImputationRules-class
      Class "ImputationRules"
```

---

**Description**

A class defining a list "rules" for imputation of SNPs. Rules are estimated population haplotype probabilities for a target SNP and one or more predictor SNPs

**Objects from the Class**

Objects are lists of *rules*. Rules are named list elements each describing imputation of a SNP by a linear regression equation. Each element is itself a list with the following elements:

**maf** The minor allele frequency of the imputed SNP

**r.squared** The squared Pearson correlation coefficient between observed and predicted SNP duration derivation of the rule.

**snp**s The names of the SNPs to be included in the regression.

**hap.probs** A numerical array containing estimated probabilities for haplotypes of the SNP to be imputed and all the predictor SNPs

If any target SNP is monomorphic, the corresponding rule is returned as NULL. An object of class `ImputationRules` has an attribute, `Max.predictors`, which gives the maximum number of predictors used for any imputation.

**Methods**

**show** `signature(object = "ImputationRules")`: prints an abbreviated listing of the rules

**summary** `signature(object = "ImputationRules")`: returns a table which shows the distribution of r-squared values achieved against the number of snps used for imputation

**plot** `signature(x="ImputationRules", y="missing")`: plots the distribution of r-squared values as a stacked bar chart

`[ ]signature(x = "ImputationRules", i = "ANY")`: subset operations

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**See Also**

[snp.imputation](#), [impute.snps](#), [single.snp.tests](#)

**Examples**

```
showClass("ImputationRules")
```

---

SingleSnpTests-class

*Classes "SingleSnpTests" and "SingleSnpTestsScore"*

---

**Description**

These are classes to hold the objects created by [single.snp.tests](#) and provide methods for extracting key elements. The class "SingleSnpTestsScore" extends class "SingleSnpTests" to include the score and score variance statistics in order to provide methods for pooling results from several studies or parts of a study

**Objects from the Class**

Objects can be created by calls of the form `new("SingleSnpTests", ...)` and `new("SingleSnpTestsScore", ...)` but, more usually, will be created by calls to [single.snp.tests](#)

**Slots**

**snp.names:** The names of the SNPs tested, as they appear as column names in the original `SnpMatrix`

**chisq:** A two-column matrix holding the 1 and 2 df association tests

**N:** The numbers of observations included in each test

**N.r2:** For tests on imputed SNPs, the product of N and the imputation  $r^2$ . Otherwise a zero-length object

**U:** (class "SingleSnpTestsScore") Score statistics

**V:** (class "SingleSnpTestsScore") Score variances

**Methods**

[ ] signature(x = "SingleSnpTests", i = "ANY"): Subsetting operator

[ ] signature(x = "SingleSnpTestsScore", i = "ANY"): Subsetting operator

**chi.squared** signature(x = "SingleSnpTests", df = "numeric"): Extract 1- and 2-df chi-squared test values

**effect.sign** signature(x = "SingleSnpTestsScore", simplify = "missing"): Extract signs of associations tested by the 1df tests

**names** signature(x="SingleSnpTests"): Extract names of test values (snp.names slot)

**p.value** signature(x = "SingleSnpTests", df = "numeric"): Evaluate 1- and 2-df test p-values

**show** signature(object = "SingleSnpTests"): List all tests and p-values

**coerce** signature(from = "SingleSnpTests", to = "data.frame"): Conversion to data frame class

**sample.size** signature(object = "SingleSnpTests"): Extract sample sizes for tests

**effective.sample.size** signature(object = "SingleSnpTests"): Extract effective sample sizes for tests. For imputed tests, these are the real sample sizes multiplied by the corresponding R-squared values for imputation

**summary** signature(object = "SingleSnpTests"): Summarize all tests and p-values

**pool2** signature(x = "SingleSnpTestsScore", y = "SingleSnpTestsScore", score = "logical"): Combine two sets of test results. Used recursively by [pool](#)

**switch.alleles** signature(x = "SingleSnpTestsScore", snps = "ANY"): Emulate, in the score vector and its (co)variances, the effect of switching of the alleles for the specified tests

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**See Also**

[single.snp.tests](#), [pool](#)

**Examples**

```
showClass("SingleSnpTests")
showClass("SingleSnpTestsScore")
```

---

SnpMatrix-class      *Class "SnpMatrix"*

---

**Description**

This class defines objects holding large arrays of single nucleotide polymorphism (SNP) genotypes generated using array technologies.

**Objects from the Class**

Objects can be created by calls of the form `new("SnpMatrix", x)` where `x` is a matrix with storage mode `"raw"`. Chips (usually corresponding to samples or subjects) define rows of the matrix while polymorphisms (loci) define columns. Rows and columns will usually have names which can be used to link the data to further data concerning samples and SNPs

**Slots**

**.Data:** Object of class `"matrix"` and storage mode `raw` Internally, missing data are coded 0 and SNP genotypes are coded 1, 2 or 3. Imputed values may not be known exactly. Such uncertain calls are grouped by probability and represented by codes 4 to 253

**Extends**

Class "matrix", from data part. Class "structure", by class "matrix". Class "array", by class "matrix". Class "vector", by class "matrix", with explicit coerce. Class "vector", by class "matrix", with explicit coerce.

**Methods**

[ ] signature(x = "SnpMatrix", i = "ANY", j = "ANY", drop = "missing"): subset operations

**cbind2** signature(x = "SnpMatrix", y = "SnpMatrix"): S4 generic function to provide cbind() for two or more matrices together by column. Row names must match and column names must not coincide. If the matrices are of the derived class [XSnpMatrix-class](#), the diploid slot values must also agree

**coerce** signature(from = "SnpMatrix", to = "numeric"): map to numeric values 0, 1, 2 or, for uncertain assignments, to the posterior expectation of the 0, 1, 2 code

**coerce** signature(from = "SnpMatrix", to = "character"): map to codes "A/A", "A/B", "B/B", ""

**coerce** signature(from = "matrix", to = "SnpMatrix"): maps numeric matrix (coded 0, 1, 2 or NA) to a SnpMatrix

**coerce** signature(from = "SnpMatrix", to = "XSnpMatrix"): maps a SnpMatrix to an XSnpMatrix. Ploidy is inferred from the genotype data since haploid genotypes should always be coded as homozygous. After inferring ploidy, heterozygous calls for haploid genotypes are set to NA

**is.na** signature(x = "SnpMatrix"): returns a logical matrix indicating whether each element is NA

**rbind2** signature(x = "SnpMatrix", y = "snp.matrix"): S4 generic function to provide rbind() for two or more matrices by row. Column names must match and duplicated row names prompt warnings

**show** signature(object = "SnpMatrix"): shows the size of the matrix (since most objects will be too large to show in full)

**summary** signature(object = "SnpMatrix"): returns summaries of the data frames returned by [row.summary](#) and [col.summary](#)

**is.na** signature(x = "SnpMatrix"): returns a logical matrix of missing call indicators

**switch.alleles** signature(x = "SnpMatrix", snps = "ANY"): Recode specified columns of the matrix to reflect allele switches

**Note**

This class requires at least version 2.3 of R

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**References**

<http://www-gene.cimr.cam.ac.uk/clayton>

**See Also**[XSnpmatrix-class](#)**Examples**

```

data(testdata)
summary(Autosomes)

# Just making it up - 3-10 will be made into NA during conversion
snps.class<-new("SnpMatrix", matrix(1:10))
snps.class
if(!isS4(snps.class)) stop("constructor is not working")

pretend.X <- as(Autosomes, 'XSnpmatrix')
if(!isS4(pretend.X)) stop("coersion to derived class is not S4")
if(class(pretend.X) != 'XSnpmatrix') stop("coersion to derived class is not working")

pretend.A <- as(Xchromosome, 'SnpMatrix')
if(!isS4(pretend.A)) stop("coersion to base class is not S4")
if(class(pretend.A) != 'SnpMatrix') stop("coersion to base class is not working")

# display the first 10 snps of the first 10 samples
print(as(Autosomes[1:10,1:10], 'character'))

# convert the empty strings (no-calls) explicitly to "NC" before
# writing to an (anonymous and temporary) csv file
csvfile <- tempfile()
write.csv(file=csvfile, gsub ('^$', 'NC',
                             as(Autosomes[1:10,1:10], 'character')
                             ), quote=FALSE)

unlink(csvfile)

```

---

XSnpmatrix-class    *Class "XSnpmatrix"*

---

**Description**

This class extends the [SnpMatrix-class](#) to deal with SNPs on the X and Y chromosomes and mitochondrial SNPs.

**Objects from the Class**

Objects can be created by calls of the form `new("XSnpmatrix", x, diploid)`. Such objects have an additional slot to objects of class "SnpMatrix" consisting of a logical array of the same length as the number of rows. This array indicates whether genotypes in that row are diploid (TRUE) or haploid (FALSE as, for example, SNPs on the X chromosome for males).

**Slots**

**.Data:** Object of class "matrix" and storage mode "raw"

**diploid:** Object of class "logical" indicating sex of samples





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chi.squared                      *Extract test statistics and p-values*

---

## Description

Generic functions to extract values from the SNP association test objects returned by various testing functions

## Usage

```
chi.squared(x, df)
deg.freedom(x)
effect.sign(x, simplify)
p.value(x, df)
sample.size(x)
effective.sample.size(x)
```

## Arguments

x	An object of class "SingleSnpTests", "SingleSnpTestsScore", or "GlmTests"
df	Either the numeric value 1 or 2 (not used when x is of class "GlmTests")
simplify	This switch is relevant when x is of class "GlmTests" and plays the same role as it does in <a href="#">sapply</a> . If simplify=TRUE, where possible the output is returned as a simple numeric vector rather than as a list

## Details

These functions operate on objects created by [single.snp.tests](#), [snp.lhs.tests](#), and [snp.lhs.tests](#).

The functions `chi.squared` and `p.value` return the chi-squared statistic and the corresponding *p*-value. The argument `df` is only used for output from `single.snp.tests`, since this function calculates both 1 df and 2 df tests for each SNP. The functions `snp.lhs.tests` and `snp.rhs.tests` potentially calculate chi-squared tests on varying degrees of freedom, which can be extracted with `deg.freedom`. The function `effect.sign` indicates the direction of associations. When applied to an output object from `snp.single.tests`, it returns +1 if the association, as measured by the 1 df test, is positive and -1 if the association is negative. Each test calculated by `GlmTests` are potentially tests of several parameters so that the effect sign can be a vector. Thus `effect.sign` returns a list of sign vectors unless, if `simplify=TRUE`, and it can be simplified as a single vector with one sign for each test. The function `sample.size` returns the number of observations actually used in the test, after exclusions due to missing data have been applied, and `effective.sample.size` returns the effective sample size which is less than the true sample size for tests on imperfectly imputed SNPs.

## Value

A numeric vector containing the chi-squared test statistics or p-values. The output vector has a `names` attribute.

**Note**

The `df` and `simplify` arguments are not always required (or legal). See above

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**See Also**

[single.snp.tests](#), [snp.lhs.tests](#), [snp.rhs.tests](#), [SingleSnpTests-class](#), [SingleSnpTestsScore-class](#), [GlmTests-class](#)

**Examples**

```
data(testdata)
tests <- single.snp.tests(cc, stratum=region, data=subject.data,
  snp.data=Autosomes, snp.subset=1:10)
chi.squared(tests, 1)
p.value(tests, 1)
```

---

`convert.snpMatrix` *Convert 'snpMatrix' objects to 'snpStats' objects*

---

**Description**

These functions convert `snpMatrix` objects to `snpStats` objects. `convert.snpMatrix` converts a single object, while `convert.snpMatrix.dir` converts all stored elements in a specified directory. They really only change the class names since most of the classes in `snpStats` are backwards-compatible with `snpMatrix`. The exception is the `ImputationRules` class; `imputation.rules` objects will need to be regenerated.

**Usage**

```
convert.snpMatrix(object)

convert.snpMatrix.dir(dir = ".", ext = "RData")
```

**Arguments**

<code>object</code>	Object to be converted
<code>dir</code>	A directory containing saved <code>snpMatrix</code> objects
<code>ext</code>	The file extension for files containing such objects

**Value**

`convert.snpMatrix` returns the converted object. `convert.snpMatrix.dir` rewrites the files in place.

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

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 example-new

*An example of intensity data for SNP genotyping*


---

### Description

The file `example-new.txt` contains some signal intensity data for testing and comparing genotype scoring algorithms

### Format

This is a text file containing data on 99 SNPs for 1550 DNA samples. One line of data appears for each SNP, starting with the SNP name and followed by 1550 pairs of intensity values. There is a header line containing variable names, with intensities labelled as `xxxxA` and `xxxxB`, where `xxxx` is the sample name.

### Details

See the package vignette "Comparing clustering algorithms".

### Source

These data were originally distributed with the "Illuminus" genotype scoring software from the Wellcome Trust Sanger Institute: <http://www.sanger.ac.uk/resources/software/illuminus/>

---

 families

*Test data for family association tests*


---

### Description

These data started life as real data derived from an affected sibling pair study of type 1 diabetes. However, original subject and SNP identifiers have been replaced by randomly chosen ones.

### Usage

```
data(families)
```

### Format

There are two objects in the loaded data file:

- `genotypes`: An object of class "`snp.matrix`" containing the SNP genotype data for both parents and affected offspring
- `pedData`: A data frame containing the standard six fields for a *LINKAGE* pedfile. The are named `familyid`, `member`, `father`, `mother`, `sex`, and `affected`

The two objects are linked by common row names.

## Details

Coding in the `pedData` frame is as in the *LINKAGE* package, except that missing data are coded NA rather than zero

## Examples

```
data(families)
summary(genotypes)
summary(pedData)
```

---

<code>filter.rules</code>	<i>Filter a set of imputation rules</i>
---------------------------	---

---

## Description

Determine which imputation rules are broken by removal of some SNPs from a study. This function is needed because, when if it emerges that genotyping of some SNPs is not reliable, necessitating their removal from study, we would also wish to remove any SNPs imputed on the basis of these unreliable SNPs.

## Usage

```
filter.rules(rules, snps.excluded, exclusions = TRUE)
```

## Arguments

<code>rules</code>	An object of class "ImputationRules" containing a set of imputation rules
<code>snps.excluded</code>	The names of the SNPs whose removal is to be investigated
<code>exclusions</code>	If TRUE, the names of the imputed SNPs which would be lost by removal of the SNPs listed in <code>snps.excluded</code> . If FALSE, the names of the imputed SNPs which would <i>not</i> be lost are returned

## Value

A character vector containing the names of imputed SNPs to be removed

## Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

## See Also

[ImputationRules-class](#), [snp.imputation](#)

## Examples

```
# No example yet
```

---

`for.exercise`*Data for exercise in use of the `snpStats` package*

---

## Description

These data have been created artificially from publicly available datasets. The SNPs have been selected from those genotyped by the International HapMap Project (<http://www.hapmap.org>) to represent the typical density found on a whole genome association chip, (the Affymetrix 500K platform, [http://www.affymetrix.com/support/technical/sample\\_data/500k\\_hapmap\\_genotype\\_data.affx](http://www.affymetrix.com/support/technical/sample_data/500k_hapmap_genotype_data.affx) for a moderately sized chromosome (chromosome 10). A study of 500 cases and 500 controls has been simulated allowing for recombination using beta software from Su and Marchini (<http://www.stats.ox.ac.uk/~marchini/software/gwas/hapgen.html>). Re-sampling of cases was weighted in such a way as to simulate three “causal” locus on this chromosome, with multiplicative effects of 1.3, 1.4 and 1.5 for each copy of the risk allele.

## Usage

```
data(for.exercise)
```

## Format

There are three data objects in the dataset:

- `snp.10`: An object of class "SnpMatrix" containing a matrix of SNP genotype calls. Rows of the matrix correspond to subjects and columns correspond to SNPs.
- `snp.support`: A conventional R data frame containing information about the SNPs typed (the chromosome position and the nucleotides corresponding to the two alleles of the SNP).
- `subject.support`: A conventional R dataframe containing information about the study subjects. There are two variables; `cc` gives case/control status (1=case), and `stratum` gives ethnicity.

## Source

The data were obtained from the diabetes and inflammation laboratory (see <http://www-gene.cimr.cam.ac.uk/todd>)

## References

<http://www-gene.cimr.cam.ac.uk/clayton>

## Examples

```
data(for.exercise)
snp.10
summary(snp.10)
summary(snp.support)
summary(subject.support)
```

---

glm.test.control    *Set up control object for GLM tests*

---

### Description

To carry out a score test for a GLM, we first fit a "base" model using the standard iteratively reweighted least squares (IRLS) algorithm and then carry out a score test for addition of further terms. This function sets various control parameters for this.

### Usage

```
glm.test.control(maxit = 20, epsilon = 1.e-5, R2Max = 0.999)
```

### Arguments

maxit	Maximum number of IRLS steps
epsilon	Convergence threshold for IRLS algorithm
R2Max	R-squared limit for aliasing of new terms

### Details

Sometimes (although not always), an iterative scheme is necessary to fit the "base" generalized linear model (GLM) before carrying out a score test for effect of adding new term(s). The `maxit` parameter sets the maximum number of iterations to be carried out, while the `epsilon` parameter sets the criterion for determining convergence. After fitting the base model, the new terms are added, but terms judged to be "aliased" are omitted. The method for determining aliasing is as follows (denoting the "design" matrix for the additional terms by  $Z$ ):

1. Step 1 Regress each column of  $Z$  on the base model matrix, using the final GLM weights from the base model fit, and replace  $Z$  with the residuals from these regressions.
2. Step 2 Consider each column of the new  $Z$  matrix in turn, regressing it on the *previous* columns (again using the weights from the base model fit). If the proportion of the weighted sum of squares "explained" by this regression exceeds `R2Max`, the term is dropped and not included in the test,

The aim of this procedure to avoid wasting degrees of freedom on columns so strongly aliased that there is little power to detect their effect.

### Value

Returns the parameters as a list in the expected order

### Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

### See Also

[snp.lhs.tests](#), [snp.rhs.tests](#)

---

`ibsCount`*Count alleles identical by state*

---

### Description

This function counts, for all pairs of subjects and across all SNPs, the total number of alleles which are identical by state (IBS)

### Usage

```
ibsCount(snp, uncertain = FALSE)
```

### Arguments

<code>snp</code>	An input object of class "SnpMatrix" or "XSnpmatrix"
<code>uncertain</code>	If FALSE, uncertain genotypes are ignored. Otherwise contributions are weighted by posterior probabilities

### Details

For each pair of subjects the function counts the total number of alleles which are IBS. For autosomal SNPs, each locus contributes 4 comparisons, since each subject carries two copies. For SNPs on the X chromosome, the number of comparisons is also 4 for female:female comparisons, but is 2 for female:male and 1 for male:male comparisons.

### Value

If there are  $N$  rows in the input matrix, the function returns an  $N*N$  matrix. The upper triangle contains the total number of comparisons and the lower triangle contains the number of these which are IBS. The diagonal contains the number of valid calls for each subject.

### Note

In genome-wide studies, the SNP data will usually be held as a series of objects (of class "SnpMatrix" or "XSnpmatrix"), one per chromosome. Note that the matrices produced by applying the `ibsCount` function to each object in turn can be added to yield the genome-wide result.

### Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

### See Also

[ibsDist](#) which calculates a distance matrix based on proportion of alleles which are IBS

### Examples

```
data(testdata)

ibs.A <- ibsCount(Autosomes[,1:100])
ibs.X <- ibsCount(Xchromosome)
```



---

ibsDist	<i>Distance matrix based on identity by state (IBS)</i>
---------	---

---

**Description**

Expresses a matrix of IBS counts (see [ibsCount](#)) as a distance matrix. The distance between two samples is returned as the proportion of allele comparisons which are *not* IBS.

**Usage**

```
ibsDist(counts)
```

**Arguments**

counts            A matrix of IBS counts as produced by the function [ibsCount](#)

**Value**

An object of class "dist" (see [dist](#))

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**See Also**

[ibsCount](#), [dist](#)

**Examples**

```
data(testdata)
ibs <- ibsCount(Xchromosome)
distance <- ibsDist(ibs)
```

---

imputation.maf	<i>Extract statistics from imputation rules</i>
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---

**Description**

These functions extract key characteristics of regression-based imputation rules stored as an object of class "ImputationRules". `imputation.maf` extracts the minor allele frequencies of the imputed SNPs and `imputation.r2` extracts the prediction  $R^2$ .

**Usage**

```
can.impute(rules)
imputation.maf(rules)
imputation.r2(rules)
imputation.nsnp(rules)
```

**Arguments**

rules            An object of class "ImputationRules"

**Details**

`can.impute` returns a logical vector identifying which rules allow a valid imputation. `imputation.maf` and `imputation.r2` extract the minor allele frequencies of the imputed SNPs and the  $R^2$  for prediction achieved when building each rule. `imputation.nsnp` returns the numbers of SNPs used in each imputation

**Value**

Either a logical vector, or a numeric vector containing the extracted values

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**See Also**

[ImputationRules-class](#), [snp.imputation](#)

**Examples**

```
# These functions are currently defined as
function (rules) sapply(rules, function(x) x$maf)
function (rules) sapply(rules, function(x) x$r2)
```

---

impute.snps

*Impute snps*

---

**Description**

Given SNPs stored in an object of class "SnpMatrix" or "XSnpMatrix" and a set of imputation rules in an object of class "ImputationRules", this function calculates imputed values.

**Usage**

```
impute.snps(rules, snps, subset = NULL, as.numeric = TRUE)
```

**Arguments**

rules            The imputation rules; an object of class "ImputationRules"

snps            The object of class "SnpMatrix" or "XSnpMatrix" containing the observed SNPs

subset          A vector describing the subset of subjects to be used. If NULL (default), then use all subjects

as.numeric      If TRUE, the output is a numeric matrix containing posterior expectations of the imputed SNPs. Otherwise the output matrix is of the same class as snps and contains uncertain genotype calls

**Value**

A matrix with imputed SNPs as columns. The imputed values are the estimated expected values of each SNP when coded 0, 1 or 2.

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**References**

Wallace, C. et al. (2010) *Nature Genetics*, **42**:68-71

**See Also**

[snp.imputation](#)

**Examples**

```
# Remove 5 SNPs from a dataset and derive imputation rules for them
data(for.exercise)
sel <- c(20, 1000, 2000, 3000, 5000)
to.impute <- snps.10[,sel]
impute.from <- snps.10[,-sel]
pos.to <- snp.support$position[sel]
pos.fr <- snp.support$position[-sel]
imp <- snp.imputation(impute.from, to.impute, pos.fr, pos.to)
# Now calculate the imputed values
imputed <- impute.snps(imp, impute.from)
```

---

 ld

---

*Pairwise linkage disequilibrium measures*


---

**Description**

This function calculates measures of linkage disequilibrium between pairs of SNPs. The two SNPs in each pair may both come from the same `SnpMatrix` object, or from two different `SnpMatrix` objects. Statistics which can be calculated are the log likelihood ratio, odds ratio, Yule's Q, covariance, D-prime, R-squared, and R.

**Usage**

```
ld(x, y = NULL, depth = NULL, stats, symmetric = FALSE)
```

**Arguments**

<code>x</code>	An object of class <code>SnpMatrix</code> or <code>XSnpMatrix</code>
<code>y</code>	(Optional) Another object of the same class as <code>x</code> . If <code>y</code> is supplied, LD statistics are calculated between each column of <code>x</code> and each column of <code>y</code> . Otherwise, they are calculated between columns of <code>x</code>
<code>depth</code>	When <code>y</code> is not supplied, this parameter is mandatory and controls the maximum lag between columns of <code>x</code> considered. Thus, LD statistics are calculated between <code>x[,i]</code> and <code>x[,j]</code> only if <code>i</code> and <code>j</code> differ by no more than <code>depth</code>

<code>stats</code>	A character vector specifying the linkage disequilibrium measures to be calculated. This should contain one or more of the strings: "LLR", "OR", "Q", "Covar", "D.prime", "R.squared", ad "R"
<code>symmetric</code>	When no <code>y</code> argument is supplied this argument controls the format of the output band matrices. If TRUE, symmetric matrices are returned and, otherwise, an upper triangular matrices are returned

### Details

For each pair of SNPs, phased haplotype frequencies are first estimated by maximum likelihood using the method described by Clayton and Leung (2007). The arrays of chosen LD statistics are then calculated and returned, either as band matrices (when `y` is not supplied), or as conventional rectangular matrices (when `y` is supplied). Band matrices are stored in compressed form as objects of class `dsCMatrix` (symmetric) or `dgCMatrix` (upper triangular). These classes are defined in the "`Matrix`" package)

### Value

If only one LD statistic is requested, the function returns either a matrix or a compressed band matrix. If more than one LD statistic is requested, a list of such objects is returned

### Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

### References

Clayton and Leung (2007) *Human Heredity*, **64**:45-51, (this paper is included in package documentation)

### See Also

"`Matrix-class`"

### Examples

```
data(testdata)
ld1 <- ld(Autosomes[, 1:50], depth=10, stats=c("D.prime", "R.squared"))
ld2 <- ld(Autosomes[, 1:20], Autosomes[, 21:25], stats="R.squared")
```

---

ld.example

*Datasets to illustrate calculation of linkage disequilibrium statistics*

---

### Description

This R data file contains data from the International HapMap project, concerning 603 SNPs spanning a one megabase region on chromosome 22, in a sample of Europeans and a sample of Africans

**Format**

There are three objects in the file:

- `ceph.lm`: A `snpMatrix` object containing the European genotype data
- `yri.lm`: A `snpMatrix` object containing the African genotype data
- `support.ld`: A dataframe containing details (chromosome position etc.

of the 603 SNPs

**Source**

<http://hapmap.ncbi.nlm.nih.gov>

**References**

The International HapMap Consortium. The International HapMap Project. *Nature* **426**:789-796 (2003)

---

<code>misinherits</code>	<i>Find non-Mendelian inheritances in family data</i>
--------------------------	---

---

**Description**

For SNP data in families, this function locates all subjects whose parents are in the dataset and tests each SNP for non-Mendelian inheritances in these trios.

**Usage**

```
misinherits(ped, id, father, mother, data = sys.parent(), snp.data)
```

**Arguments**

<code>ped</code>	Pedigree identifiers
<code>id</code>	Subject identifiers
<code>father</code>	Identifiers for subjects' fathers
<code>mother</code>	Identifiers for subjects' mothers
<code>data</code>	A data frame in which to evaluate the previous four arguments
<code>snp.data</code>	An object of class " <code>SnpMatrix</code> " containing the SNP genotypes to be tested

**Details**

The first four arguments are usually derived from a "pedfile". If a data frame is supplied for the `data` argument, the first four arguments will be evaluated in this frame. Otherwise they will be evaluated in the calling environment. If the arguments are missing, they will be assumed to be in their usual positions in the pedfile data frame i.e. in columns one to four. If the pedfile data are obtained from a dataframe, the row names of the `data` and `snp.data` files will be used to align the pedfile and SNP data. Otherwise, these vectors will be assumed to be in the same order as the rows of `snp.data`.

**Value**

A logical matrix. Rows are subjects with any non-Mendelian inheritances and columns are SNPs with any non-Mendelian inheritances. The body of the matrix details whether each subject has non-Mendelian inheritance at each SNP. If a subject has no recorded genotype for a specific SNP, the corresponding element of the output matrix is set to NA.

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**See Also**

[tdt.snp](#)

**Examples**

```
data(families)
misinherits(data=pedData, snp.data=genotypes)
```

---

mvtests

*Multivariate SNP tests*

---

**Description**

This function calculates multivariate score tests between a multivariate (or multinomial) phenotype and sets of SNPs

**Usage**

```
mvtests(phenotype, sets, stratum, data = sys.parent(), snp.data, rules = NULL, complete = FALSE, uncertain = FALSE, score = FALSE)
```

**Arguments**

phenotype	Either a factor (for a multinomial phenotype) or a matrix (for a multivariate phenotype)
sets	A list of sets of SNPs to be tested against the phenotype
stratum	(Optional) a stratifying variable
data	A data frame in which phenotype and stratum reside. If absent these are assumed to be in the parent frame and correctly aligned with the rows of snp.data
snp.data	An object of class <code>SnpMatrix</code> containing the SNP data
rules	(Optional) A set of imputation rules. The function then carries out tests on imputed SNPs
complete	If <code>TRUE</code> each test will use only subjects who have complete data for the phenotype and all SNPs in the set to be tested. If <code>FALSE</code> , then complete data for the phenotype is required, but tests are based upon complete pairs of SNPs
uncertain	If <code>TRUE</code> , uncertain genotype calls will be used in the tests (scored by their posterior expectations). Otherwise such calls are treated as missing
score	If <code>TRUE</code> , the score vectors and their variance-covariance matrices are saved in the output object for further processing

**Details**

Currently `complete=FALSE` is not implemented

**Value**

An object of class `snp.tests.glm` or `GlmTests.score` depending on whether `score` is set to `FALSE` or `TRUE` in the call

**Note**

This is an experimental version

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**Examples**

```
## No example yet
```

---

`plotUncertainty` *Plot posterior probabilities of genotype assignment*

---

**Description**

The `snpStats` package allows for storage of uncertain genotype assignments in a one byte "raw" variable. The probabilities of assignment form a three-vector, subject to the linear constraint that they sum to 1.0; their possible values are grouped into 253 different classes. This function displays counts of these classes on a two-dimensional isometric plot.

**Usage**

```
plotUncertainty(snp, nlevels = 10, color.palette = heat.colors(nlevels))
```

**Arguments**

<code>snp</code>	One or more columns of a <code>SnpMatrix</code> object
<code>nlevels</code>	Probability cells are coloured according to frequency. This argument gives the number of colours that can be used
<code>color.palette</code>	The colour palette to be used

**Details**

The plot takes the form of an equilateral triangle in which each apex represents a certain assignment to one of the three genotypes. A point within the triangle represents, by the perpendicular distance from each side, the three probabilities. Each of the 253 probability classes is represented by a hexagonal cell, coloured according to its frequency in the data, which is also written within the cell

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**Examples**

```
## No example available yet
```

---

```
pool
```

---

*Pool test results from several studies or sub-studies*

---

**Description**

Given the same set of "score" tests carried out in several studies or in several different sub-samples within a study, this function pools the evidence by summation of the score statistics and score variances. It combines tests produced by `single.snp.tests` or by `snp.lhs.tests` and `snp.rhs.tests`.

**Usage**

```
pool(..., score = FALSE)
```

**Arguments**

<code>...</code>	Objects holding the (extended) test results. These must be of class <code>SingleSnpTests.score</code> or <code>snp.tests.glm</code>
<code>score</code>	Is extended score information to be returned in the output object? Relevant only for <code>SingleSnpTestsScore</code> objects

**Details**

This function works by recursive calls to the generic function `pool2` which pools the results of two studies.

**Value**

An object of same class as the input objects (optionally without the `.score`) extension. Tests are produced for the *union* of SNPs tested in all the input objects.

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**See Also**

`pool2`, `SingleSnpTestsScore-class`, `GlmTests-class`, `single.snp.tests`, `snp.lhs.tests`, `snp.rhs.tests`

**Examples**

```
# An artificial example which simply doubles the size of a study
data(testdata)
sst <- single.snp.tests(snp.data=Autosomes, cc, data=subject.data,
  score=TRUE)
sst2 <- pool(sst, sst)
summary(sst2)
```



---

 pool2

*Pool results of tests from two independent datasets*


---

**Description**

Generic function to pool results of tests from two independent datasets. It is not designed to be called directly, but is called recursively by `pool`

**Usage**

```
pool2(x, y, score)
```

**Arguments**

<code>x, y</code>	Objects holding the (extended) test results. These must be of class <code>SingleSnpTests.score</code> or <code>snp.tests.glm</code>
<code>score</code>	Is extended score information to be returned in the output object?

**Value**

An object of same class as the input objects (optionally without the `.score`) extension. Tests are produced for the *union* of SNPs tested in all the input objects.

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**See Also**

`pool`, `SingleSnpTestsScore-class`, `GlmTests-class`, `single.snp.tests`, `snp.lhs.tests`, `snp.rhs.tests`

---

 pp

*Unpack posterior probabilities from one-byte codes*


---

**Description**

In `snpStats`, the three "posterior" probabilities corresponding to the possible values of an uncertain genotype are packed into a single byte code (with, of course, some loss in accuracy). This function, which is provided as an aid to writing new functions, unpacks the posterior probabilities from the single byte codes.

**Usage**

```
pp(x, transpose = FALSE)
```

**Arguments**

<code>x</code>	A vector, length N, which can be coerced into type <code>raw</code>
<code>transpose</code>	If <code>FALSE</code> , the result is an Nx3 matrix of posterior probabilities. If <code>TRUE</code> , a 3xN matrix is returned.

**Value**

A numeric matrix

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**Examples**

```
##
## Read imputed data from a file produced by MACH
##
path <- system.file("extdata/mach1.out.mlprob.gz", package="snpStats")
mach <- read.mach(path)
pp(mach[1:50, 10])
```

---

qq.chisq

*Quantile-quantile plot for chi-squared tests*


---

**Description**

This function plots ranked observed chi-squared test statistics against the corresponding expected order statistics. It also estimates an inflation (or deflation) factor, lambda, by the ratio of the trimmed means of observed and expected values. This is useful for inspecting the results of whole-genome association studies for overdispersion due to population substructure and other sources of bias or confounding.

**Usage**

```
qq.chisq(x, df=1, x.max, main="QQ plot",
         sub=paste("Expected distribution: chi-squared (",df," df)", sep=""),
         xlab="Expected", ylab="Observed",
         conc=c(0.025, 0.975), overdisp=FALSE, trim=0.5,
         slope.one=FALSE, slope.lambda=FALSE, pvals=FALSE,
         thin=c(0.25,50), oor.pch=24, col.shade="gray", ...)
```

**Arguments**

x	A vector of observed chi-squared test values
df	The degrees of freedom for the tests
x.max	If present, truncate the observed value (Y) axis at <code>abs(x.max)</code> . If <code>x.max</code> is negative, the y-axis will extend to <code>abs(x.max)</code> even if the observed data do not
main	The main heading
sub	The subheading
xlab	x-axis label (default "Expected")
ylab	y-axis label (default "Observed")
conc	Lower and upper probability bounds for concentration band for the plot. Set this to NA to suppress this

<code>overdisp</code>	If TRUE, an overdispersion factor, lambda, will be estimated and used in calculating concentration band
<code>trim</code>	Quantile point for trimmed mean calculations for estimation of lambda. Default is to trim at the median
<code>slope.one</code>	Is a line of slope one to be superimposed?
<code>slope.lambda</code>	Is a line of slope lambda to be superimposed?
<code>pvals</code>	Are P-values to be indicated on an axis drawn on the right-hand side of the plot?
<code>thin</code>	A pair of numbers indicating how points will be thinned before plotting (see Details). If NA, no thinning will be carried out
<code>oor.pch</code>	Observed values greater than <code>x.max</code> are plotted at <code>x.max</code> . This argument sets the plotting symbol to be used for out-of-range observations
<code>col.shade</code>	The colour with which the concentration band will be filled
<code>...</code>	Further graphical parameter settings to be passed to <code>points()</code>

### Details

To reduce plotting time and the size of plot files, the smallest observed and expected points are thinned so that only a reduced number of (approximately equally spaced) points are plotted. The precise behaviour is controlled by the parameter `thin`, whose value should be a pair of numbers. The first number must lie between 0 and 1 and sets the proportion of the X axis over which thinning is to be applied. The second number should be an integer and sets the maximum number of points to be plotted in this section.

The "concentration band" for the plot is shown in grey. This region is defined by upper and lower probability bounds for each order statistic. The default is to use the 2.5 Note that this is not a simultaneous confidence region; the probability that the plot will stray outside the band at some point exceeds 95

When required, the dispersion factor is estimated by the ratio of the observed trimmed mean to its expected value under the chi-squared assumption.

### Value

The function returns the number of tests, the number of values omitted from the plot (greater than `x.max`), and the estimated dispersion factor, lambda.

### Note

All tests must have the same number of degrees of freedom. If this is not the case, I suggest transforming to p-values and then plotting  $-2\log(p)$  as chi-squared on 2 df.

### Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

### References

Devlin, B. and Roeder, K. (1999) Genomic control for association studies. *Biometrics*, **55**:997-1004

### See Also

[single.snp.tests](#), [snp.lhs.tests](#), [snp.rhs.tests](#)

**Examples**

```
## See example the single.snp.tests() function
```

---

```
read.beagle          Read genotypes imputed by the BEAGLE program
```

---

**Description**

The BEAGLE program generates, for each SNP and each subject, posterior probabilities for the three genotypes. This function reads such data as a `SnpMatrix` object, storing the posterior probabilities to as much accuracy allowed by a one-byte coding

**Usage**

```
read.beagle(file, rownames=NULL, nsnp = NULL, header=TRUE)
```

**Arguments**

<code>file</code>	The input file name. This file may be gzipped.
<code>rownames</code>	The row names (sample identifiers) for the matrix
<code>nsnp</code>	The number of SNPs to be read in. This corresponds with the number of lines in the input file. If not supplied, the function does a preliminary pass to determine the number of lines
<code>header</code>	Set this <code>TRUE</code> if the file contains a header line (it won't for older versions of BEAGLE)

**Details**

In later versions of BEAGLE, row names are listed on a header line. However, if the `rownames` argument is supplied, this will take precedence over the header line. If there is no header line and no row names are supplied, names are generated as `Sample1`, `Sample2` etc.

No provision is made for data for the X chromosome. Such data must be first read as a `SnpMatrix` and subsequently coerced to an `XSnpMatrix` object

**Value**

an object of class `SnpMatrix`

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**See Also**

[SnpMatrix-class](#)

**Examples**

```
##---- No example available yet
```

---

read.impute	<i>Read genotypes imputed by the IMPUTE2 program</i>
-------------	--

---

### Description

The IMPUTE2 program generates, for each SNP and each subject, posterior probabilities for the three genotypes. This function reads such data as a `SnpMatrix` object, storing the posterior probabilities to as much accuracy allowed by a one-byte coding

### Usage

```
read.impute(file, rownames = NULL, nsnp = NULL, snpcol = 2)
```

### Arguments

<code>file</code>	The input file name. This file may be gzipped.
<code>rownames</code>	The row names for the output object. Note that these correspond to groups of three columns in the input file. If not supplied, names are generated as <code>Sample1, Sample2</code> etc.
<code>nsnp</code>	The number of SNPs to be read in. This corresponds with the number of lines in the input file. If not supplied, the function does a preliminary pass to determine the number of lines
<code>snpcol</code>	Which column of the input will be used as the SNP name. Default is column 2

### Details

No provision is made for data for the X chromosome. Such data must be first read as a `SnpMatrix` and subsequently coerced to an `XSnpMatrix` object

### Value

an object of class `SnpMatrix`

### Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

### See Also

[SnpMatrix-class](#)

### Examples

```
##---- No example available yet
```

---

read.long

*Read SNP genotype data in long format*


---

### Description

This function reads SNP genotype data from a file in which each line refers to a single genotype call. Replaces the earlier function `read.snps.long`.

### Usage

```
read.long(file, samples, snps,
          fields = c(snp = 1, sample = 2, genotype = 3, confidence = 4,
                    allele.A = NA, allele.B = NA),
          split = "\\t| +", gcodes, no.call = "", threshold = NULL,
          lex.order = FALSE, verbose = FALSE)
```

### Arguments

<code>file</code>	Name(s) of file(s) to be read (can be gzipped)
<code>samples</code>	Either a vector of sample identifiers, or the number of samples to be read. If a single file is to be read and this argument is omitted, the file will be scanned initially and all samples will be included
<code>snps</code>	Either a vector of SNP identifiers, or the number of SNPs to be read. If a single file is to be read and this argument is omitted, the file will be scanned initially and all SNPs will be included
<code>fields</code>	A named vector giving the locations of the required fields. See Details below
<code>split</code>	A regular expression specifying how the input line will be split into fields. The default value specifies separation of fields by a TAB character, or by one or more blanks
<code>gcodes</code>	When the genotype is read as a single field, this argument specifies how it is handled. See Details below.
<code>no.call</code>	The string which indicates "no call" for either a genotype or (when the genotype is read as two allele fields) an allele
<code>threshold</code>	A vector of length 2 giving the lower and higher acceptable limits for the confidence score
<code>lex.order</code>	If <code>TRUE</code> , the alleles at each locus will be in lexicographical order. Otherwise, ordering of alleles is arbitrary, depending on the order in which they are encountered
<code>verbose</code>	If <code>TRUE</code> , this turns on output from the function. Otherwise only error and warning messages are produced

### Details

Each line on the input file represents a single call and is split into fields using the function `strsplit`. The required fields are extracted according to the `fields` argument. This *must* contain the locations of the sample and snp identifier fields and *either* the location of a genotype field *or* the locations of two allele fields.

If the `samples` and `snps` arguments contain vectors of character strings, a `SnpMatrix` is created with these row and column names and the genotype values are "cherry-picked" from the input file. If either, or both, of these arguments are specified simply as numbers, then these numbers determine the *dimensions* of the `SnpMatrix` created. In this case `samples` and/or `SNPs` are included in the `SnpMatrix` on a first-come-first-served basis. If either or both of these arguments are omitted, a preliminary scan of the input file is carried out to find the missing sample and/or SNP identifiers. In this scan, when a sample or SNP identifier differs from that in the previous line, but is identical to one previously found, then all the relevant identifiers are assumed to have been found. This implies that the file must be sorted, in some consistent order, by sample and by SNP (although either one of these may vary fastest).

If the genotype is read as a single field, its handling is specified by the `gcodes` argument. If this is absent, `NULL`, or `NA`, then the genotype is assumed to be represented by a two-character field (the two characters representing the two alleles). If `gcodes` is a single string, then this is assumed to be a regular expression which will split the genotype field into two allele fields. Otherwise, `gcode` must be an array of length three, specifying the three genotype codes in the order "AA", "AB", "BB"

### Value

If the genotype is read as a single field matching one of three specified codes, the function returns an object of class `SnpMatrix`. Otherwise it returns a list whose first element is the `SnpMatrix` object and whose second element is a dataframe containing the allele codes, with the SNP identifiers as row names. Note that allele codes only occur in this file if they occur in a genotype which was accepted. Thus, monomorphic SNPs have `allele.B` coded as `NA`, and SNPs which never pass confidence score filters have both alleles coded as `NA`.

### Note

Unlike `read.snps.long`, this function is written entirely in R and may not be particularly fast. However, it imposes no restrictions on the allele codes recognized.

Homozygous genotypes are assumed to be represented in the input file by coding both alleles to the same value. No special provision is made to read `XSnpMatrix` objects; such data should first be read as a `SnpMatrix` and then coerced to an `XSnpMatrix` using `new` or `as`.

### Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

### See Also

[SnpMatrix-class](#), [XSnpMatrix-class](#)

### Examples

```
##  
## No example supplied yet  
##
```

---

read.mach	<i>Read genotypes imputed by the MACH program</i>
-----------	---

---

### Description

This routine reads imputed genotypes generated by the MACH program. With the `--mle` and `--mldetails` options in force this program generates a `.mlprob` output file which contains probabilities of assignments. These are stored as uncertain genotype calls in a `SnpMatrix` object

### Usage

```
read.mach(file, colnames = NULL, nrow = NULL)
```

### Arguments

<code>file</code>	The name of the <code>.mlprob</code> file. This may be gzipped
<code>colnames</code>	The column names. If absent, names are generated as <code>SNP1</code> , <code>SNP2</code> , etc.
<code>nrow</code>	If known the number of rows of data on the file. If not supplied, it is determined by a preliminary pass through the data

### Details

No routine is explicitly available for data on chromosome X. Such data should first be read as a `SnpMatrix` and then coerced to an `XSnpMatrix` object

### Value

An object of class `SnpMatrix`

### Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

### See Also

[SnpMatrix-class](#)

### Examples

```
##---- No example available yet
```



---

read.pedfile                    *Read a pedfile as "SnpMatrix" object*

---

### Description

Reads diallelic data in linkage "pedfile" format, with one line of data per sample (subject) containing six mandatory fields followed by pairs of fields, one pair for each locus, giving the two alleles observed.

### Usage

```
read.pedfile(file, n, snps, which, split = "\t| +", sep = ".", na.strings = "0",
```

### Arguments

<code>file</code>	The input pedfile. This may be (but need not be) gzipped
<code>n</code>	(Optional) The number of lines of data to be read. If not supplied the pedfile is read once and rewound to determine how many lines it contains
<code>snps</code>	(Optional) Either a character vector giving the names of the loci, or a single character variable giving the name of a locus information file from which these can be read. This file is assumed to be white-space delimited with one line per locus and no header line. If this argument is not supplied, locus names are generated as a numerical sequence, prefixed by <code>locus</code> and a separator character
<code>which</code>	(Optional) If locus names are to be read from a file, this argument should specify which column contains the names. If not supplied, the first column giving unique locus names is used
<code>split</code>	A "regexp" specifying how the input pedfile will be split into fields. The default value specifies either a TAB character or one or more spaces
<code>sep</code>	The separator character used in constructing row and column names of the output <code>SnpMatrix</code> object
<code>na.strings</code>	One or more strings to be set to NA. Any field taking one of these values will be set to NA
<code>lex.order</code>	If TRUE, then alleles will be allocated to internal 1 and 2 values in lexicographic order. Otherwise they are converted in the order in which they are encountered when reading the file (the default setting)

### Details

Row names for the output `SnpMatrix` object and for the accompanying subject description dataframe are taken as the pedigree identifiers, when these provide the required unique identifiers. When these are duplicated, an attempt is made to use the pedigree-member identifiers instead but, when these too are duplicated, row names are obtained by concatenating, with a separator character, the pedigree and pedigree-member identifiers.

### Value

A list, comprising

genotypes	The output genotype data as an object of class "SnpMatrix". If either the pedigree or pedigree-member identifiers in the ped file are not duplicated, these are used for the row names of the output object. Otherwise these two fields are concatenated, separated by sep
fam	A dataframe containing the first six fields in the pedfile. The row names will correspond with those of the SnpMatrix
map	A dataframe giving the alleles at each locus. If locus names were obtained from a dataframe read from an existing file, then the allele information is simply appended to this frame. Otherwise a new dataframe is created. The row names will correspond with the column names of the SnpMatrix

### Note

This function is written entirely in R and may not be particularly fast. However, it imposes no restrictions on the allele codes recognized.

Homozygous genotypes may be represented in the input file either (a) by coding both alleles to the same value, or (b) setting the second allele to "missing" (as specified by the `missing.allele` argument). No special provision is made to read `XSnpMatrix` objects; such data should first be read as a `SnpMatrix` and then coerced to an `XSnpMatrix` using `new` or `as`.

### Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

### See Also

[SnpMatrix-class](#), [XSnpMatrix-class](#)

### Examples

```
##
## No example supplied yet
##
```

---

read.plink

*Read a PLINK binary data file as a SnpMatrix*

---

### Description

The package PLINK saves genome-wide association data in groups of three files, with the extensions `.bed`, `.bim`, and `.fam`. This function reads these files and creates an object of class "SnpMatrix"

### Usage

```
read.plink(bed, bim, fam, na.strings = c("0", "-9"), sep = "." , select.subjects
```

**Arguments**

<code>bed</code>	The name of the file containing the packed binary SNP genotype data. It should have the extension <code>.bed</code> ; if it doesn't, then this extension will be appended
<code>bim</code>	The file containing the SNP descriptions
<code>fam</code>	The file containing subject (and, possibly, family) identifiers. This is basically a tab-delimited "pedfile"
<code>na.strings</code>	Strings in <code>.bam</code> and <code>.fam</code> files to be recoded as NA
<code>sep</code>	A separator character for constructing unique subject identifiers
<code>select.subjects</code>	A numeric vector indicating a subset of subjects to be selected from the input file (see details)
<code>select.snps</code>	Either a numeric or a character vector indicating a subset of SNPs to be selected from the input file (see details)

**Details**

If the `bed` argument does not contain a filename with the file extension `.bed`, then this extension is appended to the argument. The remaining two arguments are optional; their default values are obtained by replacing the `.bed` filename extension by `.bim` and `.fam` respectively. See the PLINK documentation for the detailed specification of these files.

The `select.subjects` or `select.snps` argument can be used to read a subset of the data. Use of `select.snps` requires that the `.bed` file is in SNP-major order (the default in PLINK). Likewise, use of `select.subjects` requires that the `.bed` file is in individual-major order. Subjects are selected by their numeric order in the PLINK files, while SNPs are selected either by order or by name. Note that the order of selected SNPs/subjects in the output objects will be the same as their order in the PLINK files.

Row names for the output `SnpMatrix` object and for the accompanying subject description dataframe are taken as the pedigree identifiers, when these provide the required unique identifiers. When these are duplicated, an attempt is made to use the pedigree-member identifiers instead but, when these too are duplicated, row names are obtained by concatenating, with a separator character, the pedigree and pedigree-member identifiers.

**Value**

A list with three elements:

<code>genotypes</code>	The output genotype data as an object of class <code>"SnpMatrix"</code> .
<code>fam</code>	A dataframe corresponding to the <code>.fam</code> file, containing the first six fields in a standard pedfile. The row names will correspond with those of the <code>SnpMatrix</code>
<code>map</code>	A dataframe corresponding to the <code>.bim</code> file. The row names correspond with the column names of the <code>SnpMatrix</code>

**Note**

No special provision is made to read `XSnpMatrix` objects; such data should first be read as a `SnpMatrix` and then coerced to an `XSnpMatrix` using `new` or `as`.

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**References**

PLINK: Whole genome association analysis toolset. <http://pngu.mgh.harvard.edu/~purcell/plink/>

**See Also**

[write.plink](#), [SnpMatrix-class](#), [XSnpMatrix-class](#)

---

read.snps.long      *Read SNP data in long format (deprecated)*

---

**Description**

Reads SNP data when organized in free format as one call per line. Other than the one call per line requirement, there is considerable flexibility. Multiple input files can be read, the input fields can be in any order on the line, and irrelevant fields can be skipped. The samples and SNPs to be read must be pre-specified, and define rows and columns of an output object of class "SnpMatrix". This function has been replaced in versions 1.3 and later by the more flexible function `read.long`.

**Usage**

```
read.snps.long(files, sample.id = NULL, snp.id = NULL, diploid = NULL,
               fields = c(sample = 1, snp = 2, genotype = 3, confidence = 4),
               codes = c("0", "1", "2"), threshold = 0.9, lower = TRUE,
               sep = " ", comment = "#", skip = 0, simplify = c(FALSE, FALSE),
               verbose = FALSE, in.order=TRUE, every = 1000)
```

**Arguments**

<code>files</code>	A character vector giving the names of the input files
<code>sample.id</code>	A character vector giving the identifiers of the samples to be read
<code>snp.id</code>	A character vector giving the names of the SNPs to be read
<code>diploid</code>	A logical array of the same length as <code>sample.id</code> , required if reading data into an <code>XSnpMatrix</code> rather than a <code>SnpMatrix</code> . This vector gives the expected ploidy for each row. If the same value suffices for all rows, then a scalar may be supplied
<code>fields</code>	A integer vector with named elements specifying the positions of the required fields in the input record. The fields are identified by the names <code>sample</code> and <code>snp</code> for the sample and SNP identifier fields, <code>confidence</code> for a call confidence score (if present) and either <code>genotype</code> if genotype calls occur as a single field, or <code>allele1</code> and <code>allele2</code> if the two alleles are coded in different fields
<code>codes</code>	Either the single string "nucleotide" denoting that coding in terms of nucleotides (A, C, G or T, case insensitive), or a character vector giving genotype or allele codes (see below)

threshold	A numerical value for the calling threshold on the confidence score
lower	If TRUE, then <code>threshold</code> represents a lower bound. Otherwise it is an upper bound
sep	The delimiting character separating fields in the input record
comment	A character denoting that any remaining input on a line is to be ignored
skip	An integer value specifying how many lines are to be skipped at the beginning of each data file
simplify	If TRUE, sample and SNP identifying strings will be shortened by removal of any common leading or trailing sequences when they are used as row and column names of the output <code>Snpmatrix</code>
verbose	If TRUE, a progress report is generated as every <code>every</code> lines of data are read
in.order	If TRUE, input lines are assumed to be in the correct order (see details)
every	See <code>verbose</code>

### Details

If nucleotide coding is not used, the `codes` argument should be a character array giving the valid codes. For genotype coding of autosomal SNPs, this should be an array of length 3 giving the codes for the three genotypes, in the order homozygous(AA), heterozygous(AB), homozygous(BB). All other codes will be treated as "no call". The default codes are "0", "1", "2". For X SNPs, males are assumed to be coded as homozygous, unless an additional two codes are supplied (representing the AY and BY genotypes). For allele coding, the `codes` array should be of length 2 and should specify the codes for the two alleles. Again, any other code is treated as "missing" and, for X SNPs, males should be coded either as homozygous or by omission of the second allele.

For nucleotide coding, nucleotides are assigned to the nominal alleles in alphabetic order. Thus, for a SNP with either "T" and "A" nucleotides in the variant position, the nominal genotypes AA, AB and BB will refer to A/A, A/T and T/T.

Although the function allows for reading into an object of class `XSnpmatrix` directly, it is usually preferable to read such data as a `Snpmatrix` (i.e. as autosomal) and to coerce it to an object of type `XSnpmatrix` later using `as(..., "X.Snpmatrix")` or `new("XSnpmatrix", ..., diploid=...)`. If `diploid` is coded NA for any subject the latter course *must* be followed, since NAs are not accepted in the `diploid` argument.

If the `in.order` argument is set TRUE, then the vectors `sample.id` and `snp.id` must be in the same order as they vary on the input file(s) and this ordering must be consistent. However, there is no requirement that either SNP or sample should vary fastest as this is detected from the input. If `in.order` is FALSE, then no assumptions about the ordering of the input file are assumed and SNP and sample identifiers are looked up in hash tables as they are read. This option must be expected, therefore, to be somewhat slower. Each file may represent a separate sample or SNP, in which case the appropriate `.id` argument can be omitted; row or column names are then taken from the file names.

### Value

An object of class `Snpmatrix` or `XSnpmatrix`.

### Note

The function will read gzipped files.

If `in.order` is TRUE, every combination of sample and snp listed in the `sample.id` and `snp.id` arguments *must* be present in the input file(s). Otherwise the function will search for any missing observation until reaching the end of the data, ignoring everything else on the way.

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**See Also**

[read.plink](#), [SnpMatrix-class](#), [XSnpMatrix-class](#)

---

row.summary	<i>Summarize rows or columns of a snp matrix</i>
-------------	--

---

**Description**

This function calculates summary statistics of each row or column of call rates and heterozygosity for each row of an object of class "SnpMatrix" or "XSnpMatrix"

**Usage**

```
row.summary(object)
col.summary(object, rules = NULL, uncertain = TRUE)
```

**Arguments**

object	genotype data as a <a href="#">SnpMatrix-class</a> or <a href="#">XSnpMatrix-class</a> object
rules	An object of class "ImputationRules". If supplied, the rules coded in this object are used, together with the snp genotype data in object, to generate imputed SNPs. The column summary of these imputed data are then returned
uncertain	If TRUE uncertain genotypes are used in calculation of allele and genotype frequencies (by scoring as posterior expectations). Otherwise, and for Hardy-Weinberg tests, they are ignored

**Value**

row.summary	returns a data frame with rows corresponding to rows of the input object and with columns/elements: <ul style="list-style-type: none"> <li>• Call.rate: Proportion of SNPs called</li> <li>• Certain.calls: Proportion of called SNPs with certain calls</li> <li>• Heterozygosity: Proportion of called SNPs which are heterozygous</li> </ul> Uncertain calls are ignored for calculating the heterozygosity.
col.summary	returns a data frame with rows corresponding to columns of the input object and with columns/elements: <ul style="list-style-type: none"> <li>• Calls: The number of valid calls</li> <li>• Call.rate: The proportion of genotypes called</li> <li>• Certain.calls: Proportion of called SNPs with certain calls</li> <li>• RAF: The "risk" allele (allele B) frequency</li> <li>• MAF: The minor allele frequency</li> <li>• P.AA: The frequency of homozygous genotype 1 (A/A)</li> <li>• P.AB: The frequency of heterozygous genotype 2 (A/B)</li> <li>• P.BB: The frequency of homozygous genotype 3 (B/B)</li> </ul>

- z.HWE: A z-test for Hardy-Weinberg equilibrium

For objects of class "X`SnpMatrix`", the following additional columns are returned:

- P.AY: The frequency of allele A in males
- P.BY: The frequency of allele B in males
- Calls.female: The number of valid calls in females (only these calls are used in the z-test for HWE)

### Note

The current version of `row.summary` does not deal with the X chromosome differently, so that males are counted as homozygous.

### Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

### Examples

```
data(testdata)
rs <- row.summary(Autosomes)
summary(rs)
cs <- col.summary(Autosomes)
summary(cs)
cs <- col.summary(Xchromosome)
summary(cs)
```

---

sample.ped.gz

*Sample datasets to illustrate data input*

---

### Description

The first five files concern data on 20 diallelic loci on 120 subjects. These data are distributed with the Haploview package (Barrett et al., 2003). The sixth file contains an additional dataset of 18 SNPs in 100 subjects, coded in long format. These six files are used in the data input vignette. The final file is a sample imputed genotype dataset distributed with the MACH imputation package, and used in the imputation vignette.

These files are stored in the `extdata` relative to the package base. Full file names can be obtained using the `system.file` function.

### Format

There are five files:

- `sample.ped.gz`: A gzipped pedfile
- `sample.info`: An accompanying locus information file
- `sample.bed`: The corresponding PLINK `.bed` file
- `sample.bim`: The PLINK `.bim` file
- `sample.fam`: The PLINK `.fam` file
- `sample-long.gz`: A sample of long-formatted data

- mach1.out.mlprob.gz: An mlprob output file from the MACH genotype imputation program. This file contains, for each imputed genotype call, posterior probabilities for the three possible genotypes

### Source

<http://www.broadinstitute.org/scientific-community/science/programs/medical-and-population-genetics/haploview/downloads> <http://www.sph.umich.edu/csg/abecasis/MACH/download>

### References

Barrett JC, Fry B, Maller J, Daly MJ.(2005) Haploview: analysis and visualization of LD and haplotype maps. *Bioinformatics*, 2005 Jan 15, [PubMed ID: 15297300]

---

single.snp.tests     *1-df and 2-df tests for genetic associations with SNPs (or imputed SNPs)*

---

### Description

This function carries out tests for association between phenotype and a series of single nucleotide polymorphisms (SNPs), within strata defined by a possibly confounding factor. SNPs are considered one at a time and both 1-df and 2-df tests are calculated. For a binary phenotype, the 1-df test is the Cochran-Armitage test (or, when stratified, the Mantel-extension test). The function will also calculate the same tests for SNPs imputed by regression analysis.

### Usage

```
single.snp.tests(phenotype, stratum, data = sys.parent(), snp.data,
  rules=NULL, subset, snp.subset, uncertain = FALSE, score=FALSE)
```

### Arguments

phenotype	A vector containing the values of the phenotype
stratum	Optionally, a factor defining strata for the analysis
data	A dataframe containing the phenotype and stratum data. The row names of this are linked with the row names of the snps argument to establish correspondence of phenotype and genotype data. If this argument is not supplied, phenotype and stratum are evaluated in the calling environment and should be in the same order as rows of snps
snp.data	An object of class "SnpMatrix" containing the SNP genotypes to be tested
rules	An object of class "ImputationRules". If supplied, the rules coded in this object are used, together with snp.data, to calculate tests for imputed SNPs
subset	A vector or expression describing the subset of subjects to be used in the analysis. This is evaluated in the same environment as the phenotype and stratum arguments
snp.subset	A vector describing the subset of SNPs to be considered. Default action is to test all SNPs in snp.data or, in imputation mode, as specified by rules



uncertain	If TRUE, uncertain genotypes are handled by replacing score contributions by their posterior expectations. Otherwise they are treated as missing. Setting this option automatically invokes use of <code>robust</code> variance estimates
score	If TRUE, the output object will contain, for each SNP, the score vector and its variance-covariance matrix

### Details

Formally, the test statistics are score tests for generalized linear models with canonical link. That is, they are inner products between genotype indicators and the deviations of phenotypes from their stratum means. Variances (and covariances) are those of the permutation distribution obtained by randomly permuting phenotype within stratum.

When the function is used to calculate tests for imputed SNPs, the test is still a score test. The score statistics are calculated from the expected value, given observed SNPs, of the score statistic if the SNP to be tested were itself observed.

The `subset` argument can either be a logical vector of length equal to the length of the vector of phenotypes, an integer vector specifying positions in the data frame, or a character vector containing names of the selected rows in the data frame. Similarly, the `snp.subset` argument can be a logical, integer, or character vector.

### Value

An object of class "SingleSnpTests". If `score` is set to TRUE, the output object will be of the extended class "SingleSnpTestsScore" containing additional slots holding the score statistics and their variances (and covariances). This allows meta-analysis using the `pool` function.

### Note

The 1 df imputation tests are described by Chapman et al. (2008) and the 2 df imputation tests are a simple extension of these. The behaviour of this function for objects of class `XSnpMatrix` is as described by Clayton (2008). Males are treated as homozygous females and corrected variance estimates are used.

### Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

### References

Chapman J.M., Cooper J.D., Todd J.A. and Clayton D.G. (2003) *Human Heredity*, **56**:18-31.  
Clayton (2008) Testing for association on the X chromosome *Biostatistics*, **9**:593-600.)

### See Also

[snp.lhs.tests](#), [snp.rhs.tests](#), [impute.snps](#), [ImputationRules-class](#), [pool](#), [SingleSnpTests-class](#), [SingleSnpTestsScore-class](#)

### Examples

```
data(testdata)
results <- single.snp.tests(cc, stratum=region, data=subject.data,
  snp.data=Autosomes, snp.subset=1:10)
print(summary(results))
```

```
# writing to an (anonymous and temporary) csv file
csvfile <- tempfile()
write.csv(file=csvfile, as(results, 'data.frame'))
unlink(csvfile)
# QQ plot
qq.chisq(chi.squared(results, 1), 1)
qq.chisq(chi.squared(results, 2), 2)
```

---

sm.compare

*Compare two SnpMatrix objects*


---

### Description

For quality control purposes, it is sometimes necessary to compare genotype data derived from different sources. This function facilitates this.

### Usage

```
sm.compare(obj1, obj2, row.wise = TRUE, col.wise = TRUE)
```

### Arguments

obj1	The first of the two <code>SnpMatrix</code> objects to be compared
obj2	The second <code>SnpMatrix</code> object
row.wise	Calculate comparison statistics aggregated in a row-wise manner
col.wise	Calculate column-wise comparison statistics

### Details

Initially row and column names of the two objects are compared to identify subsets of subjects and SNPs which they have in common. Then, every instance of a SNP genotype in the two objects are compared and agreements and disagreements counted by row and/or by column.

### Value

If only one of the row-wise and column-wise summaries are to be calculated, the return value is a matrix with rows defined by subjects or SNPs and columns giving counts of:

Agree	Agreements (all)
Disagree	Disagreements (all)
NA.agree	Genotype coded NA in both objects
NA.disagree	Genotype coded NA in only one object
Hom.agree	Objects agree and genotype is homozygous
Hom.switch	Genotype coded as homozygous in both objects, but alleles switched
Het.agree	Genotype coded as heterozygous in both objects
Het.Hom	Genotype coded as heterozygous in one object and homozygous in the other

If both row-wise and column-wise summaries are computed (the default behaviour), the function returns a list containing two matrices of the form described above. These are named `row.wise` and `col.wise`

**Note**

No special provision is yet made for objects of class `XSnpmatrix`, in which haploid calls are coded as homozygous.

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**See Also**

[SnpMatrix-class](#), [XSnpmatrix-class](#)

**Examples**

```
##  
## No example yet available  
##
```

---

snp.cbind

*Bind together two or more SnpMatrix objects*

---

**Description**

These functions bind together two or more objects of class `"SnpMatrix"` or `"XSnpmatrix"`.

**Usage**

```
# cbind(...)  
# rbind(...)  
snp.cbind(...)  
snp.rbind(...)
```

**Arguments**

...                    Objects of class `"SnpMatrix"` or `"XSnpmatrix"`.

**Details**

These functions reproduce the action of the standard functions `cbind` and `rbind`. These are constrained to work by recursive calls to the generic functions `cbind2` and `rbind2` which take just two arguments. This is somewhat inefficient in both time and memory use when binding more than two objects, so the functions `snp.cbind` and `snp.rbind`, which take multiple arguments, are also supplied.

When matrices are bound together by column, row names must be identical, column names must not be duplicated and, for objects of class `XSnpmatrix` the contents of the `Female` slot must match. When matrices are bound by row, column names must be identical and duplications of row names generate warnings.

**Value**

A new matrix, of the same type as the input matrices.

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**See Also**

[cbind](#), [rbind](#)

**Examples**

```
data(testdata)
# subsetting ( Autosomes[c(1:9,11:19,21:29),] ) is quicker. this is just for illustrating
# rbind and cbind
first <- Autosomes[1:9,]
second <- Autosomes[11:19,]
third <- Autosomes[21:29,]
result1 <- rbind(first, second, third)
result2 <- snp.rbind(first, second, third)
all.equal(result1, result2)

result3 <- Autosomes[c(1:9,11:19,21:29),]
all.equal(result1, result3)

first <- Autosomes[,1:9]
second <- Autosomes[,11:19]
third <- Autosomes[,21:29]
result1 <- cbind(first, second, third)
result2 <- snp.cbind(first, second, third)
all.equal(result1, result2)

result3 <- Autosomes[,c(1:9,11:19,21:29)]
all.equal(result1, result3)

first <- Xchromosome[1:9,]
second <- Xchromosome[11:19,]
third <- Xchromosome[21:29,]
result1 <- rbind(first, second, third)
result2 <- snp.rbind(first, second, third)
all.equal(result1, result2)

result3 <- Xchromosome[c(1:9,11:19,21:29),]
all.equal(result1, result3)

first <- Xchromosome[,1:9]
second <- Xchromosome[,11:19]
third <- Xchromosome[,21:29]
result1 <- cbind(first, second, third)
result2 <- snp.cbind(first, second, third)
all.equal(result1, result2)

result3 <- Xchromosome[,c(1:9,11:19,21:29)]
all.equal(result1, result3)
```

---

`snp.cor`*Correlations with columns of a SnpMatrix*

---

### Description

This function calculates Pearson correlation coefficients between columns of a `SnpMatrix` and columns of an ordinary matrix. The two matrices must have the same number of rows. All valid pairs are used in the computation of each correlation coefficient.

### Usage

```
snp.cor(x, y, uncertain = FALSE)
```

### Arguments

<code>x</code>	An $N$ by $M$ <code>SnpMatrix</code>
<code>y</code>	An $N$ by $P$ general matrix
<code>uncertain</code>	If <code>TRUE</code> , uncertain genotypes are replaced by posterior expectations. Otherwise these are treated as missing values

### Details

This can be used together with `xxt` and `eigen` to calculate standardized loadings in the principal components

### Value

An  $M$  by  $P$  matrix of correlation coefficients

### Note

This version cannot handle X chromosomes

### Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

### See Also

[xxt](#)

### Examples

```
# make a SnpMatrix with a small number of rows
data(testdata)
small <- Autosomes[1:100,]
# Calculate the X.X-transpose matrix
xx <- xxt(small, correct.for.missing=TRUE)
# Calculate the principal components
pc <- eigen(xx, symmetric=TRUE)$vectors
# Calculate the loadings in first 10 components */
loadings <- snp.cor(small, pc[,1:10])
```

---

snp.imputation      *Calculate imputation rules*

---

### Description

Given two set of SNPs typed in the same subjects, this function calculates rules which can be used to impute one set from the other in a subsequent sample. The function can also calculate rules for imputing each SNP in a single dataset from other SNPs in the same dataset

### Usage

```
snp.imputation(X, Y, pos.X, pos.Y, phase=FALSE, try=50, stopping=c(0.95, 4, 0.05),
               use.hap=c(1.0, 0.0), em.cntnl=c(50,0.01,10,0.01), minA=5)
```

### Arguments

X	An object of class "SnpMatrix" or "XSnpMatrix" containing observations of the SNPs to be used for imputation ("predictor SNPs")
Y	An object of same class as X containing observations of the SNPs to be imputed in a future sample ("target SNPs"). If this argument is missing, then target SNPs are also drawn from X
pos.X	The positions of the predictor SNPs. Can be missing if there is no Y argument and the columns of X are in genome position order
pos.Y	The positions of the target SNPs. Only required when a Y argument is present
phase	See "Details" below
try	The number of potential predictor SNPs to be considered in the stepwise regression procedure around each target SNP. The nearest try predictor SNPs to each target SNP will be considered
stopping	Parameters of the stopping rule for the stepwise regression (see below)
use.hap	Parameters to control use of the haplotype imputation method (see below)
em.cntnl	Parameters to control test for convergence of EM algorithm for fitting phased haplotypes (see below)
minA	A minimum data quantity measure for estimating pairwise linkage disequilibrium (see below)

### Details

The routine first carries out a series of step-wise least-square regression analyses in which each Y SNP is regressed on the nearest try predictor (X) SNPs. If phase is TRUE, the regressions will be calculated at the chromosome (haplotype) level, variances being simply  $p(1-p)$  and covariances estimated from the estimated two-locus haplotypes (this option is not yet implemented). Otherwise, the analysis is carried out at the genotype level based on conventional variance and covariance estimates using the "pairwise.complete.obs" missing value treatment (see cov). New SNPs are added to the regression until either (a) the value of  $R^2$  exceeds the first parameter of stopping, (b) the number of "tag" SNPs has reached the maximum set in the second parameter of stopping, or (c) the change in  $R^2$  does not achieve the target set by the third parameter of stopping. If the third parameter of stopping is NA, this last test is replaced by a test for improvement in the Akaike information criterion (AIC).

After choosing the set of "tag" SNPs in this way, a prediction rule is generated either by calculating phased haplotype frequencies, either (a) under a log-linear model for linkage disequilibrium with only first order association terms fitted, or (b) under the "saturated" model. These methods do not differ if there is only one tag SNP but, otherwise, choice between methods is controlled by the `use.hap` parameters. If the prediction, as measure by  $R^2$  achieved with the log-linear smoothing model exceeds a threshold (the first parameter of `use.hap`) then this method is used. Otherwise, if the gain in  $R^2$  achieved by using the second method exceeds the second parameter of `use.hap`, then the second method is used. Current experience is that, the log-linear method is rarely preferred with reasonable choices for `use.hap`, and imputation is much faster when the second method only is considered. The current default ensures that this second method is used, but the other possibility might be considered if imputing from very small samples; however this code is not extensively tested and should be regarded as experimental.

The argument `em.ctrl` controls convergence testing for the EM algorithm for fitting haplotype frequencies and the IPF algorithm for fitting the log-linear model. The first parameter is the maximum number of EM iterations, and the second parameter is the threshold for the change in log likelihood below which the iteration is judged to have converged. The third and fourth parameters give the maximum number of IPF iterations and the convergence tolerance. There should be no need to change the default values.

All SNPs selected for imputation must have sufficient data for estimating pairwise linkage disequilibrium with each other and with the target SNP. The statistic chosen is based on the four-fold tables of two-locus haplotype frequencies. If the frequencies in such a table are labelled  $a, b, c$  and  $d$  then, if  $ad > bc$  then  $t = \min(a, d)$  and, otherwise,  $t = \min(b, c)$ . The cell frequencies  $t$  must exceed `minA` for all pairwise comparisons.

### Value

An object of class "ImputationRules".

### Note

The `phase=TRUE` option is not yet implemented

### Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

### References

- Chapman J.M., Cooper J.D., Todd J.A. and Clayton D.G. (2003) *Human Heredity*, **56**:18-31.  
Wallace, C. et al. (2010) *Nature Genetics*, **42**:68-71

### See Also

[ImputationRules-class](#), [imputation.maf](#), [imputation.r2](#)

### Examples

```
# Remove 5 SNPs from a dataset and derive imputation rules for them
data(for.exercise)
sel <- c(20, 1000, 2000, 3000, 5000)
to.impute <- snps.10[,sel]
impute.from <- snps.10[,-sel]
pos.to <- snp.support$position[sel]
```

```
pos.fr <- snp.support$position[~sel]
imp <- snp.imputation(impute.from, to.impute, pos.fr, pos.to)
```

---

snp.lhs.estimates *Logistic regression with SNP genotypes as dependent variable*

---

## Description

Under the assumption of Hardy-Weinberg equilibrium, a SNP genotype is a binomial variate with two trials for an autosomal SNP or with one or two trials (depending on sex) for a SNP on the X chromosome. With each SNP in an input "SnpMatrix" as dependent variable, this function fits a logistic regression model. The Hardy-Weinberg assumption can be relaxed by use of a "robust" option.

## Usage

```
snp.lhs.estimates(snp.data, base.formula, add.formula, subset, snp.subset,
                 data = sys.parent(), robust = FALSE, uncertain = FALSE,
                 control=glm.test.control(maxit=20, epsilon=1.e-5, R2Max=0.999))
```

## Arguments

snp.data	The SNP data, as an object of class "SnpMatrix" or "XSnpMatrix"
base.formula	A formula object describing a base model containing those terms which are to be fitted but for which parameter estimates are not required (the dependent variable is omitted from the model formula)
add.formula	A formula object describing the additional terms in the model for which parameter estimates are required (again, the dependent variable is omitted)
subset	An array describing the subset of observations to be considered
snp.subset	An array describing the subset of SNPs to be considered. Default action is to test all SNPs.
data	The data frame in which base.formula, add.formula and subset are to be evaluated
robust	If TRUE, Hardy-Weinberg equilibrium will is not assumed in calculating the variance-covariance matrix of parameter estimates
uncertain	If TRUE, uncertain genotypes are used and scored by their posterior expectations. Otherwise they are treated as missing. If set, this option forces robust variance estimates
control	An object giving parameters for the IRLS algorithm fitting of the base model and for the acceptable aliasing amongst new terms to be tested. See <a href="#">glm.test.control</a>

## Details

The model fitted is the union of the base.formula and add.formula models, although parameter estimates (and their variance-covariance matrix) are only generated for the parameters of the latter. The "robust" option causes a Huber-White "sandwich" estimate of the variance-covariance matrix to be used in place of the usual inverse second derivative matrix of the log-likelihood (which assumes Hardy-Weinberg equilibrium). If a data argument is supplied, the snp.data and data objects are aligned by rowname. Otherwise all variables in the model formulae are assumed to be stored in the same order as the columns of the snp.data object.



**Value**

An object of class `GlmEstimates`

**Note**

A factor (or several factors) may be included as arguments to the function `strata(...)` in the `base.formula`. This fits all interactions of the factors so included, but leads to faster computation than fitting these in the normal way. Additionally, a `cluster(...)` call may be included in the base model formula. This identifies clusters of potentially correlated observations (e.g. for members of the same family); in this case, an appropriate robust estimate of the variance-covariance matrix of parameter estimates is calculated.

If uncertain genotypes (e.g. as a result of imputation) are used, the interpretation of the regression coefficients is questionable.

A known bug is that the function fails when no `data` argument is supplied and the base model formula contains no variables (`~1`). A work-round is to create a data frame to hold the variables in the models and pass this as `data=`.

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**See Also**

[GlmEstimates-class](#), [snp.lhs.tests](#)

**Examples**

```
data(testdata)
test1 <-
snp.lhs.estimates(Autosomes[,1:10], ~cc, ~region, data=subject.data)
test2 <-
snp.lhs.estimates(Autosomes[,1:10], ~strata(region), ~cc,
  data=subject.data)
test3 <-
snp.lhs.estimates(Autosomes[,1:10], ~cc, ~region, data=subject.data, robust=TRUE)
test4 <-
snp.lhs.estimates(Autosomes[,1:10], ~strata(region), ~cc,
  data=subject.data, robust=TRUE)
test5 <- snp.lhs.estimates(Autosomes[,1:10], ~region+sex, ~cc, data=subject.data, robust=
print(test1)
print(test2)
print(test3)
print(test4)
print(test5)
```

## Description

Under the assumption of Hardy-Weinberg equilibrium, a SNP genotype is a binomial variate with two trials for an autosomal SNP or with one or two trials (depending on sex) for a SNP on the X chromosome. With each SNP in an input "SnpMatrix" as dependent variable, this function first fits a "base" logistic regression model and then carries out a score test for the addition of further term(s). The Hardy-Weinberg assumption can be relaxed by use of a "robust" option.

## Usage

```
snp.lhs.tests(snp.data, base.formula, add.formula, subset, snp.subset,
              data = sys.parent(), robust = FALSE, uncertain = FALSE,
              control=glm.test.control(maxit=20, epsilon=1.e-5, R2Max=0.999),
              score=FALSE)
```

## Arguments

snp.data	The SNP data, as an object of class "SnpMatrix" or "XSnpMatrix"
base.formula	A formula object describing the base model, with dependent variable omitted
add.formula	A formula object describing the additional terms to be tested, also with dependent variable omitted
subset	An array describing the subset of observations to be considered
snp.subset	An array describing the subset of SNPs to be considered. Default action is to test all SNPs.
data	The data frame in which base.formula, add.formula and subset are to be evaluated
robust	If TRUE, a test which does not assume Hardy-Weinberg equilibrium will be used
uncertain	If TRUE, uncertain genotypes are used and scored by their posterior expectations. Otherwise they are treated as missing. If set, this option forces robust variance estimates
control	An object giving parameters for the IRLS algorithm fitting of the base model and for the acceptable aliasing amongst new terms to be tested. See <a href="#">glm.test.control</a>
score	Is extended score information to be returned?

## Details

The tests used are asymptotic chi-squared tests based on the vector of first and second derivatives of the log-likelihood with respect to the parameters of the additional model. The "robust" form is a generalized score test in the sense discussed by Boos(1992). If a data argument is supplied, the snp.data and data objects are aligned by rowname. Otherwise all variables in the model formulae are assumed to be stored in the same order as the columns of the snp.data object.

## Value

An object of class [snp.tests.glm](#) or [GlmTests.score](#) depending on whether score is set to FALSE or TRUE in the call.

**Note**

A factor (or several factors) may be included as arguments to the function `strata(...)` in the `base.formula`. This fits all interactions of the factors so included, but leads to faster computation than fitting these in the normal way. Additionally, a `cluster(...)` call may be included in the base model formula. This identifies clusters of potentially correlated observations (e.g. for members of the same family); in this case, an appropriate robust estimate of the variance of the score test is used.

A known bug is that the function fails when no `data` argument is supplied and the base model formula contains no variables ( $\sim 1$ ). A work-round is to create a data frame to hold the variables in the models and pass this as `data=`.

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**References**

Boos, Dennis D. (1992) On generalized score tests. *The American Statistician*, **46**:327-333.

**See Also**

[GlmTests-class](#), [GlmTestsScore-class](#), [glm.test.control](#), [snp.rhs.tests](#) [single.snp.tests](#)  
[SnpMatrix-class](#), [XSnpMatrix-class](#)

**Examples**

```
data(testdata)
snp.lhs.tests(Autosomes[,1:10], ~cc, ~region, data=subject.data)
snp.lhs.tests(Autosomes[,1:10], ~strata(region), ~cc,
  data=subject.data)
```

---

`snp.pre.multiply`    *Pre- or post-multiply a SnpMatrix object by a general matrix*

---

**Description**

These functions first standardize the input `SnpMatrix` in the same way as does the function `xxt`. The standardized matrix is then either pre-multiplied (`snp.pre.multiply`) or post-multiplied (`snp.post.multiply`) by a general matrix. Allele frequencies for standardizing the input `SnpMatrix` may be supplied but, otherwise, are calculated from the input `SnpMatrix`

**Usage**

```
snp.pre.multiply(snps, mat, frequency=NULL, uncertain = FALSE)
snp.post.multiply(snps, mat, frequency=NULL, uncertain = FALSE)
```

**Arguments**

snp	An object of class "SnpMatrix" or "XSnpMatrix"
mat	A general (numeric) matrix
frequency	A numeric vector giving the allele (relative) frequencies to be used for standardizing the columns of snps. If NULL, allele frequencies will be calculated internally. Frequencies should refer to the second (B) allele
uncertain	If TRUE, uncertain genotypes are replaced by posterior expectations. Otherwise these are treated as missing values

**Details**

The two matrices must be conformant, as with standard matrix multiplication. The main use envisaged for these functions is the calculation of factor loadings in principal component analyses of large scale SNP data, and the application of these loadings to other datasets. The use of externally supplied allele frequencies for standardizing the input SnpMatrix is required when applying loadings calculated from one dataset to a different dataset

**Value**

The resulting matrix product

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**See Also**

[xxt](#)

**Examples**

```
##--
##-- Calculate first two principal components and their loading, and verify
##--
# Make a SnpMatrix with a small number of rows
data(testdata)
small <- Autosomes[1:20,]
# Calculate the X.X-transpose matrix
xx <- xxt(small, correct.for.missing=FALSE)
# Calculate the first two principal components and corresponding eigenvalues
eigvv <- eigen(xx, symmetric=TRUE)
pc <- eigvv$vector[,1:2]
ev <- eigvv$value[1:2]
# Calculate loadings for first two principal components
Dinv <- diag(1/sqrt(ev))
loadings <- snp.pre.multiply(small, Dinv %*% t(pc))
# Now apply loadings back to recalculate the principal components
pc.again <- snp.post.multiply(small, t(loadings) %*% Dinv)
print(cbind(pc, pc.again))
```

---

snp.rhs.estimates *Fit GLMs with SNP genotypes as independent variable(s)*

---

## Description

This function fits a generalized linear model with phenotype as dependent variable and with a series of SNPs (or small sets of SNPs) as predictor variables. Optionally, one or more potential confounders of a phenotype-genotype association may be included in the model. In order to protect against misspecification of the variance function, "robust" estimates of the variance-covariance matrix of estimates may be calculated in place of the usual model-based estimates.

## Usage

```
snp.rhs.estimates(formula, family = "binomial", link, weights, subset, data
= parent.frame(), snp.data,
  rules = NULL, sets = NULL, robust = FALSE, uncertain = FALSE,
  control=glm.test.control(maxit=20, epsilon=1.e-5, R2Max=0.999))
```

## Arguments

formula	The model formula, with phenotype as dependent variable and any potential confounders as independent variables. Note that parameter estimates are not returned for these model terms
family	A string defining the generalized linear model family. This currently should (partially) match one of "binomial", "Poisson", "Gaussian" or "gamma" (case-insensitive)
link	A string defining the link function for the GLM. This currently should (partially) match one of "logit", "log", "identity" or "inverse". The default action is to use the "canonical" link for the family selected
data	The dataframe in which the model formula is to be interpreted
snp.data	An object of class "SnpMatrix" or "XSnpMatrix" containing the SNP data
rules	Optionally, an object of class "ImputationRules"
sets	Either a vector of SNP names (or numbers) for the SNPs to be added to the model formula, or a logical vector of length equal to the number of columns in snp.data or a list of short vectors defining sets of SNPs to be included (see Details)
weights	"Prior" weights in the generalized linear model
subset	Array defining the subset of rows of data to use
robust	If TRUE, robust tests will be carried out
uncertain	If TRUE, uncertain genotypes are used and scored by their posterior expectations. Otherwise they are treated as missing
control	An object giving parameters for the IRLS algorithm fitting of the base model and for the acceptable aliasing amongst new terms to be tested. See <a href="#">glm.test.control</a>

## Details

Homozygous SNP genotypes are coded 0 or 2 and heterozygous genotypes are coded 1. For SNPs on the X chromosome, males are coded as homozygous females. For X SNPs, it will often be appropriate to include sex of subject in the base model (this is not done automatically). The "robust" option causes Huber-White estimates of the variance-covariance matrix of the parameter estimates to be returned. These protect against mis-specification of the variance function in the GLM, for example if binary or count data are overdispersed,

If a `data` argument is supplied, the `snp.data` and `data` objects are aligned by rowname. Otherwise all variables in the model formulae are assumed to be stored in the same order as the columns of the `snp.data` object.

Usually SNPs to be fitted in models will be referenced by name. However, they can also be referenced by number, indicating the appropriate column in the input `snp.data`. They can also be referenced by a logical selection vector of length equal to the number of columns in `snp.data`.

If the `rules` argument is supplied, SNPs may be imputed using these rules and included in the model.

## Value

An object of class `GlmEstimates`

## Note

A factor (or several factors) may be included as arguments to the function `strata(...)` in the formula. This fits all interactions of the factors so included, but leads to faster computation than fitting these in the normal way. Additionally, a `cluster(...)` call may be included in the base model formula. This identifies clusters of potentially correlated observations (e.g. for members of the same family); in this case, an appropriate robust estimate of the variance of the parameter estimates is used.

If uncertain genotypes (e.g. as a result of imputation) are used, the interpretation of the regression coefficients is questionable; the regression coefficient for an imperfectly measurement of a variable is not a biased (attenuated) estimate of the coefficient of the variable measured.

## Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

## See Also

[GlmEstimates-class](#), [snp.lhs.estimate](#), [snp.rhs.test](#), [SnpMatrix-class](#), [XSnpMatrix-class](#)

## Examples

```
data(testdata)
test <- snp.rhs.estimate(cc~strata(region), family="binomial",
  data=subject.data, snp.data= Autosomes, sets=1:10)
print(test)
test2 <- snp.rhs.estimate(cc~region+sex, family="binomial",
  data=subject.data, snp.data= Autosomes, sets=1:10)
print(test2)
test.robust <- snp.rhs.estimate(cc~strata(region), family="binomial",
  data=subject.data, snp.data= Autosomes, sets=1:10, robust=TRUE)
print(test.robust)
```

---

snp.rhs.tests      *Score tests with SNP genotypes as independent variable*

---

### Description

This function fits a generalized linear model with phenotype as dependent variable and, optionally, one or more potential confounders of a phenotype-genotype association as independent variable. A series of SNPs (or small groups of SNPs) are then tested for additional association with phenotype. In order to protect against misspecification of the variance function, "robust" tests may be selected.

### Usage

```
snp.rhs.tests(formula, family = "binomial", link, weights, subset, data = parent
  snp.data, rules=NULL, tests=NULL, robust = FALSE, uncertain=FALSE,
  control=glm.test.control(maxit=20, epsilon=1.e-5, R2Max=0.999),
  allow.missing=0.01, score=FALSE)
```

### Arguments

formula	The base model formula, with phenotype as dependent variable
family	A string defining the generalized linear model family. This currently should (partially) match one of "binomial", "Poisson", "Gaussian" or "gamma" (case-insensitive)
link	A string defining the link function for the GLM. This currently should (partially) match one of "logit", "log", "identity" or "inverse". The default action is to use the "canonical" link for the family selected
data	The dataframe in which the base model is to be fitted
snp.data	An object of class "SnpMatrix" or "XSnpMatrix" containing the SNP data
rules	An object of class "ImputationRules". If supplied, the rules coded in this object are used, together with snp.data, to calculate tests for imputed SNPs
tests	Either a vector of SNP names (or numbers) for the SNPs to be tested, or a logical vector of length equal to the number of columns in snp.data, or a list of short numeric or character vectors defining groups of SNPs to be tested (see Details)
weights	"Prior" weights in the generalized linear model
subset	Array defining the subset of rows of data to use
robust	If TRUE, robust tests will be carried out
uncertain	If TRUE, uncertain genotypes are used and scored by their posterior expectations. Otherwise they are treated as missing
control	An object giving parameters for the IRLS algorithm fitting of the base model and for the acceptable aliasing amongst new terms to be tested. See <a href="#">glm.test.control</a>
allow.missing	The maximum proportion of SNP genotype that can be missing before it becomes necessary to refit the base model
score	Is extended score information to be returned?

## Details

The tests used are asymptotic chi-squared tests based on the vector of first and second derivatives of the log-likelihood with respect to the parameters of the additional model. The "robust" form is a generalized score test in the sense discussed by Boos(1992). The "base" model is first fitted, and a score test is performed for addition of one or more SNP genotypes to the model. Homozygous SNP genotypes are coded 0 or 2 and heterozygous genotypes are coded 1. For SNPs on the X chromosome, males are coded as homozygous females. For X SNPs, it will often be appropriate to include sex of subject in the base model (this is not done automatically).

If a `data` argument is supplied, the `snp.data` and `data` objects are aligned by rowname. Otherwise all variables in the model formulae are assumed to be stored in the same order as the columns of the `snp.data` object.

Usually SNPs to be used in tests will be referenced by name. However, they can also be referenced by number, a positive number indicating the appropriate column in the input `snp.data`, and a negative number indicating (minus) a position in the `rules` list. They can also be referenced by a logical selection vector of length equal to the number of columns in `snp.data`. Sets of tests involving more than one SNP are referenced by a list and can use a mixture of observed and imputed SNPs. If the `tests` argument is missing, single SNP tests are carried out; if a `rules` is given, all *imputed* SNP tests are calculated, otherwise all SNPs in the input `snp.data` matrix are tested. But note that, for single SNP tests, the function `single.snp.tests` will often achieve the same result much faster.

## Value

An object of class `GlmTests` or `GlmTestsScore` depending on whether `score` is set to `FALSE` or `TRUE` in the call.

## Note

A factor (or several factors) may be included as arguments to the function `strata(...)` in the formula. This fits all interactions of the factors so included, but leads to faster computation than fitting these in the normal way. Additionally, a `cluster(...)` call may be included in the base model formula. This identifies clusters of potentially correlated observations (e.g. for members of the same family); in this case, an appropriate robust estimate of the variance of the score test is used.

## Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

## References

Boos, Dennis D. (1992) On generalized score tests. *The American Statistician*, **46**:327-333.

## See Also

[GlmTests-class](#), [GlmTestsScore-class](#), [single.snp.tests](#), [snp.lhs.tests](#), [impute.snps](#), [ImputationRules-class](#), [SnpMatrix-class](#), [XSnpMatrix-class](#)

## Examples

```
data(testdata)
slt3 <- snp.rhs.tests(cc~strata(region), family="binomial",
  data=subject.data, snp.data= Autosomes, tests=1:10)
```



```
print(slt3)
```

---

snpStats-package    *SnpMatrix and XSnpmatrix classes and methods*

---

## Description

Classes and statistical methods for large SNP association studies, extending the snpMatrix package

## Details

Package: snpStats  
 Version: 1.4.1  
 Date: 2011-12-19  
 Depends: R(>= 2.3.0), survival, methods, Matrix  
 Imports: graphics, grDevices, methods, stats, survival, utils, Matrix  
 Suggests: hexbin  
 License: GPL-3  
 URL: <http://www-gene.cimr.cam.ac.uk/clayton>  
 Collate: ss.R contingency.table.R convert.R compare.R glm-test.R imputation.R indata.R long.R misc.R ld.R mvtests.R  
 LazyLoad: yes  
 biocViews: Microarray, SNP, GeneticVariability  
 Packaged: 2011-12-19  
 Built: R 2.13.1; x86\_64-unknown-linux-gnu; 2011-12-19 15:21:42 UTC; unix

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Fst	Calculate fixation indices
GlmEstimates-class	Class "GlmEstimates"
GlmTests-class	Classes "GlmTests" and "GlmTestsScore"
ImputationRules-class	Class "ImputationRules"
SingleSnpTests-class	Classes "SingleSnpTests" and "SingleSnpTestsScore"
SnpMatrix-class	Class "SnpMatrix"
XSnpmatrix-class	Class "XSnpmatrix"
chi.squared	Extract test statistics and p-values
convert.snpMatrix	Convert 'snpMatrix' objects to 'snpStats' objects
example-new	An example of intensity data for SNP genotyping
families	Test data for family association tests
filter.rules	Filter a set of imputation rules
for.exercise	Data for exercise in use of the snpStats package
glm.test.control	Set up control object for GLM tests
ibsCount	Count alleles identical by state
ibsDist	Distance matrix based on identity by state (IBS)
imputation.maf	Extract statistics from imputation rules
impute.snps	Impute snps

ld	Pairwise linkage disequilibrium measures
ld.example	Datasets to illustrate calculation of linkage disequilibrium statistics
misinherits	Find non-Mendelian inheritances in family data
mvtests	Multivariate SNP tests
plotUncertainty	Plot posterior probabilities of genotype assignment
pool	Pool test results from several studies or sub-studies
pool2	Pool results of tests from two independent datasets
pp	Unpack posterior probabilities from one-byte codes
qq.chisq	Quantile-quantile plot for chi-squared tests
read.beagle	Read genotypes imputed by the BEAGLE program
read.impute	Read genotypes imputed by the IMPUTE2 program
read.long	Read SNP genotype data in long format
read.mach	Read genotypes imputed by the MACH program
read.pedfile	Read a pedfile as '"SnpMatrix"' object
read.plink	Read a PLINK binary data file as a SnpMatrix
read.snps.long	Read SNP data in long format (deprecated)
row.summary	Summarize rows or columns of a snp matrix
sample.info	Sample datasets to illustrate data input
single.snp.tests	1-df and 2-df tests for genetic associations with SNPs (or imputed SNPs)
sm.compare	Compare two SnpMatrix objects
snp.cbind	Bind together two or more SnpMatrix objects
snp.cor	Correlations with columns of a SnpMatrix
snp.imputation	Calculate imputation rules
snp.lhs.estimates	Logistic regression with SNP genotypes as dependent variable
snp.lhs.tests	Score tests with SNP genotypes as dependent variable
snp.pre.multiply	Pre- or post-multiply a SnpMatrix object by a general matrix
snp.rhs.estimates	Fit GLMs with SNP genotypes as independent variable(s)
snp.rhs.tests	Score tests with SNP genotypes as independent variable
snpStats-package	SnpMatrix and XSnpMatrix classes and methods
switch.alleles	Switch alleles in columns of a SnpMatrix or in test results
tdt.snp	1-df and 2-df tests for genetic associations with SNPs (or imputed SNPs) in family data
test.allele.switch	Test for switch of alleles between two collections
testdata	Test data for the snpStats package
write.SnpMatrix	Write a SnpMatrix object as a text file
write.plink	Write files for analysis in the PLINK toolset
xxt	X.X-transpose for a standardized SnpMatrix

Further information is available in the following vignettes:

data-input-vignette	Data input (source)
differences	snpMatrix-differences (source)
imputation-vignette	Imputation and meta-analysis (source)
ld-vignette	LD statistics (source)
pca-vignette	Principal components analysis (source)
snpStats-vignette	snpStats introduction (source)
tdt-vignette	TDT tests (source)

**Author(s)**

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Maintainer: David Clayton <david.clayton@cimr.cam.ac.uk>

---

switch.alleles      *Switch alleles in columns of a SnpMatrix or in test results*

---

**Description**

This is a generic function which can be applied to objects of class "SnpMatrix" or "XSnpMatrix" (which hold SNP genotype data), or to objects of class "SingleSnpTestsScore" or "GlmTests" (which hold association test results). In the former case, specified SNPs can be recoded as if the alleles were switched (so that *AA* genotypes become *BB* and vice-versa while *AB* remain unchanged). In the latter case, test results are modified *as if* alleles had been switched.

**Usage**

```
switch.alleles(x, snps)
```

**Arguments**

x	The input object, of class "SnpMatrix", "XSnpMatrix", "SingleSnpTestsScore", or "GlmTests"
snps	A vector of type integer, character or logical specifying the SNP to have its alleles switched

**Value**

An object of the same class as the input object

**Note**

Switching alleles for SNPs has no effect on test results. These functions are required when carrying out meta-analysis, bringing together several sets of results. It is then important that alleles line up in the datasets to be combined. It is often more convenient (and faster) to apply this process to the test result objects rather than to the genotype data themselves.

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**See Also**

[SnpMatrix-class](#), [XsnpMatrix-class](#), [SingleSnpTests-class](#), [GlmTests-class](#)

**Examples**

```
data(testdata)
which <- c("173774", "173811")
Asw <- switch.alleles(Autosomes, which)
col.summary(Autosomes[,which])
col.summary(Asw[,which])
```

---

tdt.snp	<i>1-df and 2-df tests for genetic associations with SNPs (or imputed SNPs) in family data</i>
---------	--

---

**Description**

Given large-scale SNP data for families comprising both parents and one or more affected offspring, this function computes 1 df tests (the TDT test) and a 2 df test based on observed and expected transmissions of genotypes. Tests based on imputation rules can also be carried out.

**Usage**

```
tdt.snp(ped, id, father, mother, affected, data = sys.parent(), snp.data,
        rules = NULL, snp.subset, check.inheritance = TRUE, robust = FALSE,
        uncertain = FALSE, score = FALSE)
```

**Arguments**

ped	Pedigree identifiers
id	Subject identifiers
father	Identifiers for subjects' fathers
mother	Identifiers for subjects' mothers
affected	Disease status (TRUE if affected, FALSE otherwise)
data	A data frame in which to evaluate the previous five arguments
snp.data	An object of class "SnpMatrix" containing the SNP genotypes to be tested
rules	An object of class "ImputationRules". If supplied, the rules coded in this object are used, together with snp.data, to calculate tests for imputed SNPs
snp.subset	A vector describing the subset of SNPs to be considered. Default action is to test all SNPs in snp.data or, in imputation mode, as specified by rules
check.inheritance	If TRUE, each affected offspring/parent trio is tested for Mendelian inheritance and excluded if the test fails. If FALSE, misinheriting trios are used but the "robust" variance option is forced
robust	If TRUE, forces the robust (Huber-White) variance option (with ped determining independent "clusters")
uncertain	If TRUE, uncertain genotypes are handed by replacing score contributions by their posterior expectations. Otherwise these are treated as missing. Setting this option automatically invokes use of robust variance estimates
score	If TRUE, the output object will contain, for each SNP, the score vector and its variance-covariance matrix

## Details

Formally, the test statistics are score tests for the "conditioning on parental genotype" (CPG) likelihood. Parametrization of associations is the same as for the population-based tests calculated by `single.snp.tests` so that results from family-based and population-based studies can be combined using `pool`.

When the function is used to calculate tests for imputed SNPs, the test is still an approximate score test. The current version does not use the family relationships in the imputation. With this option, the robust variance estimate is forced.

The first five arguments are usually derived from a "pedfile". If a data frame is supplied for the `data` argument, the first five arguments will be evaluated in this frame. Otherwise they will be evaluated in the calling environment. If the arguments are missing, they will be assumed to be in their usual positions in the pedfile data frame i.e. in columns one to four for the identifiers and column six for disease status (with affected coded 2). If the pedfile data are obtained from a dataframe, the row names of the `data` and `snp.data` files will be used to align the pedfile and SNP data. Otherwise, these vectors will be assumed to be in the same order as the rows of `snp.data`.

The `snp.subset` argument can be a logical, integer, or character vector.

If imputed rather than observed SNPs are tested, or if `check.inheritance` is set to `FALSE`, the "robust" variance estimate is used regardless of the value supplied for the `robust` argument.

## Value

An object of class `"SingleSnpTests"`. If `score=TRUE`, the output object will be of the extended class `"SingleSnpTestsScore"` containing additional slots holding the score statistics and their variances (and covariances). This allows meta-analysis using the `pool` function.

## Note

When the snps are on the X chromosome (i.e. when the `snp.data` argument is of class `"XSnpMatrix"`), the tests are constructed in the same way as was described by Clayton (2008) for population-based association tests i.e. assuming that genotype relative risks for males mirror those of homozygous females

## Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

## References

Clayton (2008) Testing for association on the X chromosome *Biostatistics*, **9**:593-600.)

## See Also

[single.snp.tests](#), [impute.snps](#), [pool](#), [ImputationRules-class](#), [SingleSnpTests-class](#), [SingleSnpTestsScore-class](#)

## Examples

```
data(families)
tdt.snp(data=pedData, snp.data=genotypes)
```

---

test.allele.switch *Test for switch of alleles between two collections*

---

### Description

When testing genotype data derived from different platforms or scoring algorithms a common problem is switching of alleles. This function provides a diagnostic for this. Input can either be two objects of class "SnpMatrix" to be examined, column by column, for allele switching, or a single "SnpMatrix" object together with an indicator vector giving group membership for its rows.

### Usage

```
test.allele.switch(snps, snps2 = NULL, split = NULL, prior.df = 1)
```

### Arguments

snps	An object of class "SnpMatrix" or "XSnpMatrix"
snps2	A second object of the same class as snps
split	If only one SnpMatrix object supplied, a vector with the same number of elements as rows of snps. It must be capable of coercion to a factor with two levels.
prior.df	A degree of freedom parameter for the prior distribution of the allele frequency prior.df (see Details)

### Details

This function calculates a Bayes factor for the comparison of the hypothesis that the alleles have been switched with the hypothesis that they have not been switched. This requires integration over the posterior distribution of the allele frequency. The prior is taken as a beta distribution with both parameters equal to prior.df so that the prior is symmetric about 0.5. The default, prior.df=1 represents a uniform prior on (0,1).

### Value

A vector containing the log (base 10) of the Bayes Factors for an allele switch.

### Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

### See Also

[SnpMatrix-class](#), [XSnpMatrix-class](#)

### Examples

```
data(testdata)
#
# Call with two SnpMatrix arguments
#
cc <- as.numeric(subject.data$cc)
lbf1 <- test.allele.switch(Autosomes[cc==1,], Autosomes[cc==2,])
```

```
#  
# Single matrix call (giving the same result)  
#  
lbf2 <- test.allele.switch(Autosomes, split=cc)
```

---

testdata

*Test data for the snpStats package*

---

## Description

This dataset comprises several data frames from a fictional (and unrealistically small) study. The dataset started off as real data from a screen of non-synonymous SNPs for association with type 1 diabetes, but the original identifiers have been removed and a random case/control status has been generated.

## Usage

```
data(testdata)
```

## Format

There are five data objects in the dataset:

- `Autosomes`: An object of class "SnpMatrix" containing genotype calls for 400 subjects at 9445 autosomal SNPs
- `Xchromosome`: An object of class "XSnpMatrix" containing genotype calls for 400 subjects at 155 SNPs on the X chromosome
- `Asnps`: A dataframe containing information about the autosomal SNPs. Here it contains only one variable, `chromosome`, indicating the chromosomes on which the SNPs are located
- `Xsnps`: A dataframe containing information about the X chromosome SNPs. Here it is empty and is only included for completeness
- `subject.data`: A dataframe containing information about the subjects from whom each row of SNP data was obtained. Here it contains:
  - `cc`: Case-control status
  - `sex`: Sex
  - `region`: Geographical region of residence

## Source

The data were obtained from the diabetes and inflammation laboratory (see <http://www-gene.cimr.cam.ac.uk/todd>)

## References

<http://www-gene.cimr.cam.ac.uk/clayton>

**Examples**

```

data(testdata)
Autosomes
Xchromosome
summary(Asnps)
summary(Xsnps)
summary(subject.data)
summary(summary(Autosomes))
summary(summary(Xchromosome))

```

---

write.plink

*Write files for analysis in the PLINK toolset*


---

**Description**

Given a `SnpMatrix` object, together with associated subject and SNP support dataframes, this function writes `.bed`, `.bim`, and `.fam` files for processing in the PLINK toolset

**Usage**

```

write.plink(file.base, snp.major = TRUE, snps,
            subject.data, pedigree, id, father, mother, sex, phenotype,
            snp.data, chromosome, genetic.distance, position, allele.1, allele.2,
            na.code = 0)

```

**Arguments**

<code>file.base</code>	A character string giving the base filename. The extensions <code>.bed</code> , <code>.bim</code> , and <code>.fam</code> are appended to this string to give the filenames of the three output files
<code>snp.major</code>	Logical variable controlling whether the <code>.bed</code> file is in SNP-major or subject-major order
<code>snps</code>	The <code>SnpMatrix</code> or <code>XSnpMatrix</code> object to be written out
<code>subject.data</code>	(Optional) A subject support dataframe. If supplied, the next six arguments (which define the fields of the PLINK <code>.fam</code> file) will be evaluated in this environment, after matching row names with the row names of <code>snps</code> . Otherwise they will be evaluated in the calling environment; they then must be of the right length and in the correct order.
<code>pedigree</code>	A pedigree (family) identifier. Default is the row names of <code>snps</code> .
<code>id</code>	An identifier of an individual within family. Default is a vector of <code>na.code</code> .
<code>father</code>	The within-family identifier of the subject's father. Default is a vector of <code>na.code</code> .
<code>mother</code>	The within-family identifier of the subject's mother. Default is a vector of <code>na.code</code> .
<code>sex</code>	Sex of the individual. Default is a vector of <code>na.code</code> . This will be coerced to type <code>numeric</code> .
<code>phenotype</code>	The primary phenotype value. Default is a vector of <code>na.code</code> . This will be coerced to type <code>numeric</code> .



snp.data	(Optional) A SNP support dataframe. If supplied, the next five arguments (which define the columns of the PLINK .bim file) will be evaluated in this environment, after matching row names with the column names of snps. Otherwise they will be evaluated in the calling environment; they then must be of the right length and in the correct order.
chromosome	The chromosome on which the SNP is located. This should either be numeric, as specified by SPLINK, or character, with "X", "Y", "XY", and "MT" for the non-numeric values. Default is a vector of na.code, or a vector of 23's if snps is a XSnpmatrix.
genetic.distance	The location of the SNP, expressed as a genetic distance. Default is a vector of na.code. This will be coerced to type numeric.
position	The physical location of the SNP, expressed in base pairs. Default is a vector of na.code. This will be coerced to type numeric.
allele.1	A character vector giving the first allele. Default is a vector of "A"s.
allele.2	A character vector giving the first allele. Default is a vector of "B"s.
na.code	The code to be written for NA in the .fam and .bin output files. It should be numeric (or capable of coercion to numeric).

### Details

For more details of required codings in .fam and .bim files, see the PLINK documentation.

### Value

Returns NULL.

### Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

### References

PLINK: Whole genome association analysis toolset. <http://pngu.mgh.harvard.edu/~purcell/plink/>

### See Also

[read.plink](#), [SnpMatrix-class](#), [XSnpmatrix-class](#)

### Examples

```
data(testdata)
## the use of as.numeric() below is not necessary since this is done
## automatically. It just illustrates use of expressions for these arguments
## Note that cc and sex are variables within the subject.data frame
## This command writes files test.bed, test.fam and test.bim
write.plink(file.base="test", snps=Autosomes,
            subject.data=subject.data, phenotype = as.numeric(cc), sex=as.numeric(sex),
            snp.major=FALSE)
```

---

write.SnpMatrix      *Write a SnpMatrix object as a text file*

---

### Description

This function is closely modelled on `write.table`. It writes an object of class `SnpMatrix` as a text file with one line for each row of the matrix. Genotypes are written in numerical form, *i.e.* as 0, 1 or 2 (where 1 denotes heterozygous) or, optionally, as a pair of alleles (each coded 1 or 2).

### Usage

```
write.SnpMatrix(x, file, as.alleles= FALSE, append = FALSE, quote = TRUE, sep =
```

### Arguments

<code>x</code>	The object to be written
<code>file</code>	The name of the output file
<code>as.alleles</code>	If TRUE, write each genotype as two alleles
<code>append</code>	If TRUE, the output is appended to the designated file. Otherwise a new file is opened
<code>quote</code>	If TRUE, row and column names will be enclosed in quotes
<code>sep</code>	The string separating entries within a line
<code>eol</code>	The string terminating each line
<code>na</code>	The string written for missing genotypes
<code>row.names</code>	If TRUE, each row will commence with the row name
<code>col.names</code>	If TRUE, the first line will contain all the column names

### Value

A numeric vector giving the dimensions of the matrix written

### Author(s)

David Clayton <david.clayton@cimr.cam.ac.uk>

### See Also

[write.table](#), [SnpMatrix-class](#), [XSnpMatrix-class](#)

xxt

*X.X-transpose for a standardized SnpMatrix***Description**

The input `SnpMatrix` is first standardized by subtracting the mean (or stratum mean) from each call and dividing by the expected standard deviation under Hardy-Weinberg equilibrium. It is then post-multiplied by its transpose. This is a preliminary step in the computation of principal components.

**Usage**

```
xxt(snp, strata = NULL, correct.for.missing = FALSE, lower.only = FALSE,
    uncertain = FALSE)
```

**Arguments**

<code>snp</code>	The input matrix, of type " <code>SnpMatrix</code> "
<code>strata</code>	A factor (or an object which can be coerced into a factor) with length equal to the number of rows of <code>snp</code> defining stratum membership
<code>correct.for.missing</code>	If <code>TRUE</code> , an attempt is made to correct for the effect of missing data by use of inverse probability weights. Otherwise, missing observations are scored zero in the standardized matrix
<code>lower.only</code>	If <code>TRUE</code> , only the lower triangle of the result is returned and the upper triangle is filled with zeros. Otherwise, the complete symmetric matrix is returned
<code>uncertain</code>	If <code>TRUE</code> , uncertain genotypes are replaced by posterior expectations. Otherwise these are treated as missing values

**Details**

This computation forms the first step of the calculation of principal components for genome-wide SNP data. As pointed out by Price et al. (2006), when the data matrix has more rows than columns it is most efficient to calculate the eigenvectors of  $X.X$ -transpose, where  $X$  is a `SnpMatrix` whose columns have been standardized to zero mean and unit variance. For autosomes, the genotypes are given codes 0, 1 or 2 after subtraction of the mean,  $2p$ , are divided by the standard deviation  $\sqrt{2p(1-p)}$  ( $p$  is the estimated allele frequency). For SNPs on the X chromosome in male subjects, genotypes are coded 0 or 2. Then the mean is still  $2p$ , but the standard deviation is  $2\sqrt{p(1-p)}$ . If the `strata` is supplied, a stratum-specific estimate value for  $p$  is used for standardization.

Missing observations present some difficulty. Price et al. (2006) recommended replacing missing observations by their means, this being equivalent to replacement by zeros in the standardized matrix. However this results in a biased estimate of the complete data result. Optionally this bias can be corrected by inverse probability weighting. We assume that the probability that any one call is missing is small, and can be predicted by a multiplicative model with row (subject) and column (locus) effects. The estimated probability of a missing value in a given row and column is then given by  $m = RC/T$ , where  $R$  is the row total number of no-calls,  $C$  is the column total of no-calls, and  $T$  is the overall total number of no-calls. Non-missing contributions to  $X.X$ -transpose are then weighted by  $w = 1/(1 - m)$  for contributions to the diagonal elements, and products of the relevant pairs of weights for contributions to off-diagonal elements.

**Value**

A square matrix containing either the complete  $X.X$ -transpose matrix, or just its lower triangle

**Warning**

The correction for missing observations can result in an output matrix which is not positive semi-definite. This should not matter in the application for which it is intended

**Note**

In genome-wide studies, the SNP data will usually be held as a series of objects (of class "SnpMatrix" or "XsnpMatrix"), one per chromosome. Note that the  $X.X$ -transpose matrices produced by applying the `xxt` function to each object in turn can be added to yield the genome-wide result.

If the matrix is converted to a correlation matrix by pre- and post-multiplying by the `sqrt` of the inverse of its diagonal, then this is an unbiased estimate of twice the kinship matrix.

**Author(s)**

David Clayton <david.clayton@cimr.cam.ac.uk>

**References**

Price et al. (2006) Principal components analysis corrects for stratification in genome-wide association studies. *Nature Genetics*, **38**:904-9

**Examples**

```
# make a SnpMatrix with a small number of rows
data(testdata)
small <- Autosomes[1:100,]
# Calculate the X.X-transpose matrix
xx <- xxt(small, correct.for.missing=TRUE)
# Calculate the principal components
pc <- eigen(xx, symmetric=TRUE)$vectors
```

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