# Package 'PCFAM'

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Title Computation of Ancestry Scores with Mixed Families and Unrelated

Type Package

Individuals

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<b>Description</b> We provide several algorithms to compute the genotype ancestry scores (such as eigenvector projections) in the case where highly correlated individuals are involved.
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PCFAM-package	Computation of ancestry scores with mixed families and unrelated in- dividuals

# **Description**

This package provides ancestry scores based on genotype data, and is robust to the presence of close-degree family members. Four main novel algorithms are represented: (i) Geometric rotation (within-family data orthogonalization); (ii) matrix substitution based on the decomposition of a target family-orthogonalized covariance matrix; (iii) covariance-preserving whitening, retaining covariances between unrelated pairs while orthogonalizing family members (Note: the function perfectwhiten generates a new dataset which keeps the same covariance structure as the original set); (iv) using family-averaged data to obtain loadings for projection of family members.

#### **Details**

Package: PCFAM
Type: Package
Version: 1.0
Date: 2016-10-11
License: GPL 2
LazyLoad: yes

# Author(s)

Yi-Hui Zhou

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## References

Computation of ancestry scores with mixed families and unrelated individuals. arXiv:1606.08416

#### **Examples**

```
X <- matrix(rbinom(1000*20,2,0.4),1000,20)
X[,1]=X[,2]*0.9
X=rowscale(X)
Xresid=residualize(X)
corXresid=cor(Xresid)
myfam=findfamilies(corXresid,0.1)
K=3
myms.pca=ms.pca(X,corXresid,0.1,K)
familyave.result=familyave(X,myfam,top=K)</pre>
```

colcenter 3

colcenter

column centering of the data matrix

# Description

This function centerizes each column of the data matrix

# Usage

colcenter(X)

# **Arguments**

Χ

input data matrix

## Value

return the data matrix with each column centered

## Author(s)

Yi-Hui Zhou

## References

Computation of ancestry scores with mixed families and unrelated individuals. Yi-Hui Zhou, J.S. Marron, Fred Wright, arXiv:1606.08416.

cov.function

Sample covariance calculator

# **Description**

Obtain a sample covariance matrix

# Usage

```
cov.function(data.matrix)
```

# Arguments

data.matrix

Input mxn data matrix

# Value

return the nxn sample covariance matrix

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## Author(s)

Yi-Hui Zhou

#### References

Computation of ancestry scores with mixed families and unrelated individuals. arXiv:1606.08416.

# **Examples**

```
X <- matrix(rbinom(1000*20,2,0.4),1000,20)
cov.X=cov.function(X)</pre>
```

familyave

Family average approach

# **Description**

This function implements the family-averaging algorithm, with loadings based on the combined data from singletons and family averages, then projected to all.

## Usage

```
familyave(Xall,myfam, top = 5)
```

# **Arguments**

Xall The original input genotype dataset

myfam The identified family IDs. Each singleton forms his/her own family.

top The number ancestry scores desired.

# **Details**

The function averages the genotype information in each family, re-inflates to have appropriate variability, andtreats as a 'singleton' for the purpose of loading calculation. Ancestry scores are obtained by projection to all.

## Value

Output the top ancestry scores by combining family data with singletons

# Author(s)

Yi-Hui Zhou

## References

fastcov 5

# **Examples**

```
X <- matrix(rbinom(1000*20,2,0.4),1000,20)
X[,1]=X[,2]*0.9
X=rowscale(X)
Xresid=residualize(X)
corXresid=cor(Xresid)
myfam=findfamilies(corXresid,0.1)
K=3
familyave.result=familyave(X,myfam,top=K)</pre>
```

fastcov

Fast covariance function

# Description

This function can generate covariance matrix faster than the regular cov() function.

# Usage

fastcov(X)

# **Arguments**

Χ

input mxn data matrix

## Value

Output nxn covariance matrix

## Note

The input data matrix has to be column scaled in advance.

# Author(s)

Yi-Hui Zhou,

# References

gr.pca

findfamilies

Find families

# Description

This function searches for pairs of individuals with high kinship based on the genotype correlation matrix.

# Usage

```
findfamilies(x, threshold = 0.4)
```

# **Arguments**

x The nxn correlation matrix of the input dataset.

threshold This threshold is used to identify close-degree relatives. Recommended values

are 0.4 to identify first-degree relatives, and 0.15 to identify first- and second-

degree relatives.

#### Value

Output numerical family ID for each individual. Individuals with the same ID are judged to be family members.

## Author(s)

Yi-Hui Zhou

# References

Computation of ancestry scores with mixed families and unrelated individuals. arXiv:1606.08416.

gr.pca

The geometric rotation approach

## **Description**

This algorithm rotates scaled genotypes among family members so that they are mutually orthogonal.

# Usage

```
gr.pca(data.input, index.family, myfam, weight, top, family.size, inflation)
```

ms.pca 7

## Arguments

data.input Input dataset, each row is for a genetic feature (SNP), each column is for indi-

vidual. Data are typically number of minor alleles, possibly imputed.

index.family Index vector to indicate the family id of each individual.

myfam This value comes directly from the output of findfamilies().

weight Weight is 0 by default. This is a deprecated weight value that can be used to

control the amount of rotation performed. A weight of zero performs full or-

thogonalization, while a weight of 1 keeps the data unchanged.

top The number of eigenvectors to be used.

family.size The number of members in each family. Used to determine rotation angles.

inflation The inflation of the data value is 0 under default. Deprecated.

#### Value

data.new The new datamatrix after the geometric rotation

topPCs The top eigenvectors topEigenvalue The top eigenvalues.

#### Author(s)

Yi-Hui Zhou

#### References

Computation of ancestry scores with mixed families and unrelated individuals. arXiv:1606.08416.

ms.pca The matrix substitution approach

#### **Description**

This function provides the matrix substitution algorithm. The main idea is to replace the high covariance value entries in the covariance matrix which are produced by family members by a small value (e.g. median covariance).

#### Usage

```
ms.pca(X, corXresid, threshold, top)
```

## **Arguments**

X The input data matrix

corXresid The correlation of the genotypes after residualization for any evidence of larger

scale ancestry. Used to identify close-degree family members in a manner robust

to large-scale ancestry.

threshold Covariance values of identified family members are set to the threshold.

top The number of ancestry scores to obtain.

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# Value

eigenvector Eigenvectors after using the matrix substitution method

myeigen The top eigenvalues and eigenvectors

## Author(s)

Yi-Hui Zhou

# References

Computation of ancestry scores with mixed families and unrelated individuals. arXiv:1606.08416.

mysqrtm

Matrix square root function

# Description

This function can find the matrix square root, without requiring a new package and often faster than other code.

# Usage

```
mysqrtm(a, symmetric = F)
```

# **Arguments**

a The input matrix

symmetric Default=FALSE. This argument indicates whether the input matrix is symmet-

ric.

# **Details**

Matrix B is said to be a square root of A if the matrix product BB is equal to A.

## Value

returns the square root matrix B

perfectwhiten 9

perfectwhiten The covariance preserving whitening function.
---

# Description

This algorithm generates a new scaled 'genotype' dataset which keeps the same covariance structure as the original data, except that family members have been made orthogonal to each other, and singletons are unchanged.

# Usage

```
perfectwhiten(Xun, Xfam, delta = 3e-04, threshold = 0.35, eta = NULL, addfuzz = F)
```

# **Arguments**

Xun	A matrix of (possibly scaled) genotypes, (number of SNPs)*(number of singletons)
Xfam	A matrix of (possibly scaled) genotypes, (number of SNPs)*(number of individuals belonging to families)
delta	A slight offset used to ensure that the target covariance matrix is of full rank
threshold	The correlation threshold used to determine pairs of relatives. The choice should be less than the degree desired. For example, 0.35 captures first degree relatives (expected correlation 0.5), 0.15 captures first and second degree relatives (expected correlation for second degree relatives is 0.25).
eta	This argument is the replacement value used for matrix substitution. The default is NULL, resulting in substitution by the median.
addfuzz	The default is FALSE. Deprecated.

# Value

	Xplusscaled	The row-scaled full genotype data, including both singletons and family members	
	Υ	The (scaled) genotype matrix after whitening, and should have a covariance matrix very close to Mtarget. Column means are zero	
Ynotcolcentered			
		The same as Y, but with column means matching those of Xplusscaled	
	М	The covariance matrix of the full data	
	Mtilde	The covariance matrix after matrix substitution of all family pairs identified with correlations exceedingeta	
	whichbig	The set of indexes of M that have correlation exceeding threshold	
	covY	The covariance matrix of Y, useful to compare to M or to Mtarget	

# Author(s)

Yi-Hui ZHou, Fred A. Wright

10 residualize

## References

Computation of ancestry scores with mixed families and unrelated individuals. arXiv:1606.08416.

## **Examples**

```
X <- matrix(rbinom(1000*20,2,0.4),1000,20)
X[,1]=X[,2]*0.9
X=rowscale(X)
Xresid=residualize(X)
library(PCFAM)
corXresid=cor(Xresid)
myfam=findfamilies(corXresid,0.1)
K=3
perfect.result=perfectwhiten(X[,which(myfam==0)],X[,which(myfam==1)])</pre>
```

residualize

Residualization and scale of the original genotype data

# **Description**

Thus function performs a simple residualization of a row-scaled genotype dataset, removing large-scale population stratification. Output is a residualized dataset appropriate for computing correlations such that family members can be easily identified. The function assumes X is row-scaled

## Usage

```
residualize(X)
```

## **Arguments**

Χ

The original input genotype dataset

#### **Details**

This function pre-treatment the data before applying the findfamily function.

# Value

Outputs the new row-scaled genotype matrix after residualization

#### Author(s)

Yi-Hui Zhou

# References

rowcol 11

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Indicators for of the row and column of the original matrix

# **Description**

This function identifies the rows and columns of elements in a matrix, e.g. the family members identified based on the correlation matrix.

# Usage

```
rowcol(I, J, elements)
```

# Arguments

J The number of columns of the matrix (scalar)

elements A vector of matrix element indexes

## Value

whichrow The rows of elements in the matrix

whichcol The columns of elements in the matrix

## Author(s)

```
Yi-Hui ZHou, Fred A. Wright
```

#### References

Computation of ancestry scores with mixed families and unrelated individuals. arXiv:1606.08416.

# **Examples**

```
X <- matrix(rbinom(1000*20,2,0.4),1000,20)
X[,1]=X[,2]*0.9
X=rowscale(X)</pre>
```

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rowscale

Scale each row of a matrix

# Description

This function scales the input matrix so that each row mean is 0 and each row (sample) variance is 1.

# Usage

rowscale(X)

# Arguments

Χ

input data matrix

# Value

Output the row-scaled matrix.

# Author(s)

Yi-Hui ZHou, Fred A. Wright

# References

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