# Package 'SmCCNet'

June 19, 2018

Type Package			
Title Sparse multiple canonical correlation network analysis tool			
Version 0.1.0			
<b>Date</b> 2018-6-15			
Author W. Jenny Shi [aut, cre], Laura Saba [aut], Katerina Kechris [aut]			
Maintainer W. Jenny Shi <wjennyshi@gmail.com></wjennyshi@gmail.com>			
<b>Description</b> A canonical correlation based framework for constructing phenotype-specific multicomics networks by integrating multiple omics data types and a quantitative phenotype of interest.			
<pre>URL https://github.com/KechrisLab/SmCCNet</pre>			
<pre>BugReports https://github.com/KechrisLab/SmCCNet/issues</pre>			
<b>Depends</b> R (>= 3.4.4)			
Imports PMA, Matrix, pbapply, igraph			
Suggests BiocStyle, knitr, rmarkdown			
VignetteBuilder knitr			
License GPL-3			
Encoding UTF-8			
LazyData true			
RoxygenNote 6.0.1			
NeedsCompilation no			
R topics documented:			
getMultiOmicsModules			
getRobustPseudoWeights			
plotMultiOmicsNetwork			
X2			
Y 7			
Index 8			

getAbar Compute the similarity matrix based on one or more canonical conlation weight vectors.	rre-
--	------

#### **Description**

Compute the similarity matrix based on the outer products of absolute canonical correlation weights.

#### Usage

```
getAbar(Ws, FeatureLabel = NULL)
```

#### **Arguments**

Ws A canonical correlation weight vector or matrix. If Ws is a matrix, then each

column corresponds to one weight vector.

FeatureLabel A  $1 \times p$  vector indicating feature names. If FeatureLabel = NULL (default), the

feature names will be indices 1 through p, where p is the total number of omics

features.

#### Value

A  $p \times p$  symmetric non-negative matrix.

#### **Examples**

```
w <- matrix(rnorm(6), nrow = 3)
Ws <- apply(w, 2, function(x)return(x/sqrt(sum(x^2))))
abar <- getAbar(Ws)</pre>
```

getMultiOmicsModules Extract multi-omics modules based on the similarity matrix.

## Description

Apply hierarchical tree cutting to the similarity matrix and extract modules that contain both omics data types.

### Usage

```
getMultiOmicsModules(Abar, P1, CutHeight = 1 - 0.1^10, PlotTree = TRUE)
```

## Arguments

Abar A similary matrix for all features (both omics data types).

P1 Total number of features for the first omics data type.

CutHeight Height threshold for the hierarchical tree cutting. Default is  $1 - 0.1^{10}$ .

PlotTree Logical. Whether to create a hierarchical tree plot.

#### Value

A list of multi-omics modules.

#### **Examples**

```
set.seed(123)
w <- rnorm(5)
w <- w/sqrt(sum(w^2))
abar <- getAbar(w)
modules <- getMultiOmicsModules(abar, P1 = 2, CutHeight = 0.5)</pre>
```

getRobustPseudoWeights

Calculate the canonical correlation weights based on sparse multiple canonical correlation analysis (SmCCA), sparse supervised canonical correlation analysis (SsCCA), or sparse canonical correlation analysis (SCCA).

#### **Description**

Integrate two omics data type (and a quantitative phenotype), and calculate the absolute canonical correlation weights for the omics features using SmCCA SsCCA, or SCCA. SmCCA and SsCCA take into account a phenotype/trait. SmCCA maximizes the total (weighted or unweighted) pairwise canonical correlation weights between two omics data types and the trait. It requires the trait to be quantitative. SsCCA prioritizes omics features based on the trait, and assigns non-zero canonical weights to features that are more correlated to the trait. SCCA does not use any trait information for computing the canonical correlation weights. All of these three methods are included in this function, along with an omic feature subsampling scheme and an optional random feature partition scheme.

### Usage

```
getRobustPseudoWeights(X1, X2, Trait, Lambda1, Lambda2, s1 = 0.7, s2 = 0.9,
NoTrait = FALSE, FilterByTrait = FALSE, SubsamplingNum = 1000,
CCcoef = NULL, trace = FALSE)
```

#### Arguments

X1	An $n \times p_1$ data matrix (e.g. mRNA) with $p_1$ features and $n$ subjects.
X2	An $n \times p_2$ data matrix (e.g. miRNA) with $p_2$ features and $n$ subjects.
Trait	An $n \times 1$ trait data matrix for the same n subjects.
Lambda1	LASSO penalty parameter for X1. Lambda1 needs to be between 0 and 1.
Lambda2	LASSO penalty parameter for X2. Lambda2 needs to be between 0 and 1.
s1	Proportion of mRNA features to be included, default at $s1 = 0.7$ . $s1$ needs to be between 0 and 1.
<b>s</b> 2	Proportion of miRNA features to be included, default at s1 = $0.9$ . s2 needs to be between 0 and 1.
NoTrait	Logical, default is FALSE. Whether trait information is provided.

FilterByTrait Logical, default is FALSE. Whether only the top (80%) features with highest

correlation to the trait will be assigned nonzero weights. The choice of 80% is

based on the PMA package.

SubsamplingNum Number of feature subsamples. Default is 1000. Larger number leads to more

accurate results, but at a higher cost.

CCcoef Optional coefficients for the SmCCA pairwise canonical correlations. If CCcoef = NULL

(default), then the objective function is the total sum of all pairwise canonical correlations. It can also be a coefficient vector that follows the column order of

combn(K, 2).

trace Logical. Whether to display the CCA algorithm trace.

#### **Details**

```
To choose SmCCA, set NoTrait = FALSE, FilterByTrait = FALSE. To choose SsCCA, set NoTrait = FALSE, FilterByTrait = TRUE. To choose SCCA, set Trait = NULL, NoTrait = TRUE.
```

#### Value

A canonical correlation weight matrix with  $p_1 + p_2$  rows. Each column is the canonical correlation weights based on subsampled X1 and X2 features. The number of columns is SubsamplineNum.

#### **Examples**

```
## For illustration, we only subsample 5 times.
set.seed(123)
# Unweighted SmCCA
W1 <- getRobustPseudoWeights(X1, X2, Trait = Y, Lambda1 = 0.05,
  Lambda2 = 0.05, s1 = 0.7, s2 = 0.9, NoTrait = FALSE, FilterByTrait = FALSE,
  SubsamplingNum = 100, CCcoef = NULL, trace = FALSE)
# Weighted SmCCA
W2 <- getRobustPseudoWeights(X1, X2, Trait = Y, Lambda1 = 0.05,
  Lambda2 = 0.05, s1 = 0.7, s2 = 0.9, NoTrait = FALSE, FilterByTrait = FALSE,
  SubsamplingNum = 100, CCcoef = c(1, 5, 5), trace = FALSE)
# SsCCA
W3 <- getRobustPseudoWeights(X1, X2, Trait = Y, Lambda1 = .05, Lambda2 = 0.5,
  s1 = 0.7, s2 = 0.9, NoTrait = FALSE, FilterByTrait = TRUE,
  SubsamplingNum = 100, CCcoef = NULL, trace = FALSE)
# SCCA
W4 <- getRobustPseudoWeights(X1, X2, Trait = NULL, Lambda1 = 0.05,
  Lambda2 = 0.05, s1 = 0.7, s2 = 0.9, NoTrait = TRUE,
  SubsamplingNum = 100, CCcoef = NULL, trace = FALSE)
```

plotMultiOmicsNetwork Plot multi-omics module networks.

#### **Description**

Plot multi-omics modules based on similarity matrix derived from pseudo canonical weights and pairwise feature correlations.

#### Usage

```
plotMultiOmicsNetwork(Abar, CorrMatrix, multiOmicsModule, ModuleIdx, P1,
    EdgeCut = 0, FeatureLabel = NULL, AddCorrSign = TRUE, SaveFile = NULL,
    ShowType1Label = TRUE, ShowType2Label = TRUE, PlotTitle = "",
    NetLayout = "lgl", ShowNodes = TRUE, VertexLabelCex = 1,
    VertexSize = 1)
```

#### **Arguments**

Abar A  $p \times p$  similarly matrix for both omics data types based on pseudo canonical

correlation weights. p is the number of total features for the two omics data

types. All entries are non-negative.

CorrMatrix A  $p \times p$  correlation matrix that provides sign information for the network.

multiOmicsModule

A list of multi-omics modules.

P1 Total number of features for the first omics data type.

EdgeCut A numerical value between 0 and 1, indicating an edge threshold for the net-

work. Any features (network nodes) without any edge strength that passes the threshold are excluded from the figure. If EdgeCut = 0 (default), then the full

module network will be created.

FeatureLabel A  $1 \times p$  vector indicating feature names. If FeatureLabel = NULL (default),

the feature names will be indices 1 through p.

AddCorrSign Logical. Whether to add a positive or negative sign to each network edge based

on pairwise feature correlations.

SaveFile A pdf file name for the figure output. If SaveFile = NULL (default), the figure

will not be saved.

ShowType1Label Logical. Whether to label the network nodes for the first omics data type.

ShowType2Label Logical. Whether to label the network nodes for the second omics data type.

PlotTitle A title for the figure. Default is without any title.

NetLayout Graphical layout for the network. Possible options are circle for circle lay-

out, sphere for 3D sphere, fr for Fruchterman-Reinhold, and 1g1 for the LGL algorithm. Refer to igraph manual for more details on the layout options.

ShowNodes Logical. Whether to show network nodes.

VertexLabelCex Scaling factor for the vertex labels.

VertexSize Size of the vertices.

6 X2

#### Value

A multi-omics network figure.

#### **Examples**

```
set.seed(123)
w <- rnorm(5)
w <- w/sqrt(sum(w^2))
abar <- getAbar(w)
modules <- getMultiOmicsModules(abar, P1 = 2, CutHeight = 0.5)
x <- cbind(X1[ ,1:2], X2[ , 1:3])
corr <- cor(x)

plotMultiOmicsNetwork(abar, corr, modules, ModuleIdx = 1, P1 = 2)</pre>
```

X1

A synthetic mRNA expression dataset.

## Description

A matrix containing simulated mRNA expression levels for 358 subjects (rows) and 500 features (columns).

## Usage

X1

#### Format

An object of class matrix with 358 rows and 500 columns.

Х2

A synthetic miRNA expression dataset.

## Description

A matrix containing simulated miRNA expression levels for 358 subjects (rows) and 100 features (columns).

## Usage

Х2

#### **Format**

An object of class matrix with 358 rows and 100 columns.

Y 7

Υ

A synthetic phenotype dataset.

## Description

A matrix containing simulated quantitative phenotype measures for 358 subjects (rows).

## Usage

Υ

## **Format**

An object of class matrix with 358 rows and 1 columns.

# **Index**

```
*Topic datasets
X1, 6
X2, 6
Y, 7

getAbar, 2
getMultiOmicsModules, 2
getRobustPseudoWeights, 3

plotMultiOmicsNetwork, 5

X1, 6
X2, 6
Y, 7
```