

# Package: SIGN (via r-universe)

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**Type** Package

**Title** Similarity Identification in Gene Expression

**Version** 0.1.0

**Date** 2018-12-02

**Description** Provides a classification framework to use expression patterns of pathways as features to identify similarity between biological samples. It provides a new measure for quantifying similarity between expression patterns of pathways.

**Depends** R(>= 3.4)

**License** GPL (>= 3)

**Imports** stats, utils, survcomp, survival, GSVA

**Encoding** UTF-8

**LazyData** true

**NeedsCompilation** yes

**biocViews** GeneExpression, Classification, Clustering, Survival

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**Repository** <https://bhaibeka.r-universe.dev>

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BubbleSort	<i>BubbleSort is a function for calculating bubble sort correlation between two vectors</i>
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## Description

BubbleSort is a function for calculating bubble sort correlation between two vectors

## Usage

```
BubbleSort(Vec1, Vec2)
```

## Arguments

Vec1	Vector of values of 1st feature across samples
Vec2	Vector of values of 2nd feature across samples

## Value

Bubble sort similarity between the two vectors

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EventRenaming	<i>EventRenaming is a function for changing annotation of censored samples to 0 and dead samples to 1 for survival analysis</i>
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**Description**

EventRenaming is a function for changing annotation of censored samples to 0 and dead samples to 1 for survival analysis

**Usage**

```
EventRenaming(EventVec, Censored_Annot)
```

**Arguments**

EventVec	Status vector for all of the samples (patients) including both samples undergone an event or censored
Censored_Annot	Index of samples censored in the dataset

**Value**

Vector of events including 0 for censoring and 1 for death

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ExpPheno_Categorize	<i>ExpPheno_Categorize is a function for grouping samples based on their survival to 3 groups of poor, good, and intermediate</i>
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**Description**

ExpPheno\_Categorize is a function for grouping samples based on their survival to 3 groups of poor, good, and intermediate

**Usage**

```
ExpPheno_Categorize(ExpMeta_List, Time_ID, Event_ID, Mad_Factor,
  MinNum_ExClass, Expression_Log2 = FALSE)
```

**Arguments**

ExpMeta_List	List containing expression matrix and metadata matrix
Time_ID	Index of time to death in metadata matrix
Event_ID	Index of event in metadata matrix
Mad_Factor	Threshold of mad in time to death values to determine poor survival group

MinNum\_ExClass Minimum number of samples that has to be kept in poor and good group (if number of samples is lower than this threshold, more samples will be added in order of survival)

Expression\_Log2 Parameter for gene expression value transformation to logarithmic scale ( $\log_2(\text{expression value}+1)$ )

**Value**

List of expression matrices, and time to event as well as event for the patients within each category of poor, intermediate or good survival

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ExpPhen_Matching	<i>ExpPhen_Matching is a function for matching samples between expression matrices and metadata matrix (clinical feature matrix)</i>
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**Description**

ExpPhen\_Matching is a function for matching samples between expression matrices and metadata matrix (clinical feature matrix)

**Usage**

```
ExpPhen_Matching(ExpMat, MetaMat, SamID_Meta)
```

**Arguments**

ExpMat	Matrix of expression of genes (samples in columns and genes in rows)
MetaMat	Matrix of clinical features (samples in columns)
SamID_Meta	Sample ID in MetaMat

**Value**

List of expression matrix and metadata of the clinical information after matching patient IDs between the expression and clinical information matrices

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ExpPhen_Subdividing	<i>ExpPhen_Subdividing is a function for grouping samples based on a clinical feature available in metadata matrix (clinical feature matrix)</i>
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**Description**

ExpPhen\_Subdividing is a function for grouping samples based on a clinical feature available in metadata matrix (clinical feature matrix)

**Usage**

```
ExpPhen_Subdividing(ExpMeta_List, SubDiv_ID)
```

**Arguments**

ExpMeta_List	List containing expression matrix and metadata matrix
SubDiv_ID	Index of the target clinical feature in metadata matrix for samples grouping

**Value**

List of expression and clinical information of patients grouped based on the specified clinical feature

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GeneMatching	<i>GeneMatching is a function to remove uncommon genes between a list of expression matrices</i>
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**Description**

GeneMatching is a function to remove uncommon genes between a list of expression matrices

**Usage**

```
GeneMatching(ExpList)
```

**Arguments**

ExpList	List of expression matrices
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**Value**

List of expression matrices restricted to the common genes between them

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Genes_SimCal	<i>Genes_SimCal is a function to calculate similarity between a set of samples and 2 reference groups of samples</i>
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**Description**

Genes\_SimCal is a function to calculate similarity between a set of samples and 2 reference groups of samples

**Usage**

```
Genes_SimCal(ExpMat_Test, ExpMat_Ref1, ExpMat_Ref2, RefIDs, TestClassIter,
             SampleIter)
```

**Arguments**

ExpMat_Test	Expression matrix for the test samples for which SIGN will identify the similarity with the 2 reference dataset
ExpMat_Ref1	Expression matrix for the 1st reference set of samples
ExpMat_Ref2	Expression matrix for the 2nd reference set of samples
RefIDs	Annotations corresponding to the 2 expression matrices (1st and 2nd names are associated with the 1st and 2nd expression matrix and )
TestClassIter	Index to be matched with RefIDs for removal of test samples from reference expression matrices
SampleIter	Index of samples in the test expression matrix exist in reference expression matrix 1 or 2

**Value**

Vector of similarity between the target samples and the 2 reference sets

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GSVA_Calculation	<i>GSVA_Calculation is a function for Calculating correlation between expression level of pathways between 2 groups using GSVA</i>
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**Description**

GSVA\_Calculation is a function for Calculating correlation between expression level of pathways between 2 groups using GSVA

**Usage**

```
GSVA_Calculation(ExpMat1, ExpMat2, GeneVec, GeneSets,
                 Name = "SampleComparison")
```

**Arguments**

ExpMat1	Expression matrix of genes in the 1st group of samples
ExpMat2	Expression matrix of genes in the 2nd group of samples
GeneVec	Name of genes in the same order as considered in ExpMat1 and ExpMat2
GeneSets	List of genes within pathways
Name	Name used for naming the columns of output matrix of correlation between the 2 groups

**Value**

Similarity of the pathway between the two expression matrices based on pearson correlation, bubble sort, and wilcoxon paaired rank test using GSEA enrichment scores of pathways

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Pathway_Grouping	<i>Pathway_Grouping is a function to make a pathway list from files containing genes within each pathway</i>
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**Description**

Pathway\_Grouping is a function to make a pathway list from files containing genes within each pathway

**Usage**

```
Pathway_Grouping(PathwayDir, Pattern)
```

**Arguments**

PathwayDir	Path of directory including the files of pathways
Pattern	Pattern should be used to select the files of pathway genes from PathwayDir

**Value**

List of genes within the pathway

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Pathway_similarity	<i>Pathway_similarity is a function for calculating correlation between expression level of pathways between 2 groups using all the available approaches in SIGN</i>
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### Description

Pathway\_similarity is a function for calculating correlation between expression level of pathways between 2 groups using all the available approaches in SIGN

### Usage

```
Pathway_similarity(ExpMat1, ExpMat2, GeneVec, GeneSets, Name)
```

### Arguments

ExpMat1	Expression matrix of genes in the 1st group of sampls
ExpMat2	Expression matrix of genes in the 2nd group of sampls
GeneVec	Name of genes in the same order as considered in ExpMat1 and ExpMat2
GeneSets	List of genes within pathways
Name	Name used for naming the columns of output matrix of correlation between the 2 groups

### Value

Similarity of the pathway between the two expression matrices using pearson correlation, bubble sort, and wilcoxon paaired rank test

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SIGN_Aggregate	<i>SIGN_Aggregate is a function to reshape the list of pathway scoring, time to death, and event and return a summary list</i>
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### Description

SIGN\_Aggregate is a function to reshape the list of pathway scoring, time to death, and event and return a summary list

### Usage

```
SIGN_Aggregate(ScoreList, TimeList, EventList)
```

### Arguments

ScoreList	List of similarity scores identified using different methodologies
TimeList	List of time to event (death) for different groups of patients
EventList	List of event vectors (death or censored) for different groups of patients



**Value**

List of scores identified for each sample as well as time to death and event of that sample

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SIGN_Ensemble_SimCal	<i>SIGN_Ensemble_SimCal is a function for Generating list fo similarities based on different pathway quantification methods and similarity measures</i>
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**Description**

SIGN\_Ensemble\_SimCal is a function for Generating list fo similarities based on different pathway quantification methods and similarity measures

**Usage**

```
SIGN_Ensemble_SimCal(ExpList, RefClassID, TestClassID, GeneID, PathwaySets)
```

**Arguments**

ExpList	List of expression matrices for different groups of samples used in the centroid classification scheme
RefClassID	Names of the matrices in the ExpList
TestClassID	ID of a matrix in ExpList to be used as test set
GeneID	Parameter to determine if gene annotations are provided as Symbols or EntrezIDs
PathwaySets	List of pathways containing gene annotations for each pathways

**Value**

List of similarities identified in both gene and pathway level

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Similarities_Wrapper	<i>Similarities_Wrapper is wrapper to identify similarities between the expression of genes in target sample and the reference expression matrix</i>
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**Description**

Similarities\_Wrapper is wrapper to identify similarities between the expression of genes in target sample and the reference expression matrix

**Usage**

```
Similarities_Wrapper(ExpMat_Test, ExpMat_Ref, GeneVec, PathwaySet, RefID,
  TestClassIter, SampleIter)
```

**Arguments**

ExpMat_Test	Expression matrix of test samples
ExpMat_Ref	Expression matrix of reference samples
GeneVec	Vector of gene names
PathwaySet	List of pathways containing gene annotations for each pathways
RefID	Class of the reference set
TestClassIter	Class of the test set (if it is the same as reference set, the target test sample will be removed fro the reference set)
SampleIter	Target test sample in ExpMat_Testto be used for comparison with ExpMat_Ref

**Value**

List of similarities between the target sample and the expression matrix of reference samples

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SimSummary_2Class	<i>SimSummary_2Class is a function to calculating similarity between two set of samples</i>
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**Description**

SimSummary\_2Class is a function to calculating similarity between two set of samples

**Usage**

```
SimSummary_2Class(SimMat1, SimMat2)
```

**Arguments**

SimMat1	Matrix of similarity of the target samples with the 1st reference matrix
SimMat2	Matrix of similarity of the target samples with the 2nd reference matrix

**Value**

Matrix of similarities of samples

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SurvivalStat\_PostProcess

*SurvivalStat\_PostProcess is a function to Extract summary statistics of the built cox model*

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### Description

SurvivalStat\_PostProcess is a function to Extract summary statistics of the built cox model

### Usage

```
SurvivalStat_PostProcess(StatList)
```

### Arguments

StatList            Summary lists of the cox models built using all the

### Value

A list including Cindex, Cindex\_std and LogTest\_pval

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Survival\_Stats

*Survival\_Stats is a function for building cox model using all the features and each feature as a separate model*

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### Description

Survival\_Stats is a function for building cox model using all the features and each feature as a separate model

### Usage

```
Survival_Stats(ScoreMat, TimeVec, EventVec)
```

### Arguments

ScoreMat            Matrix of feature values used for survival prediction  
 TimeVec            Vectore of time to death of samples (patients)  
 EventVec            Vector of events for the samples (patients) as being dead or censored

### Value

A list containing summary of a cox model using all of the features and separate cox models for each feature

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TSC                      *TSC is a function to calculate transcriptional similarity coefficient between two biological pathways*

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**Description**

TSC is a function to calculate transcriptional similarity coefficient between two biological pathways

**Usage**

```
TSC(PathwayExp1, PathwayExp2)
```

**Arguments**

PathwayExp1      Expression matrix of genes within the chosen pathway in the 1st set of samples  
PathwayExp2      Expression matrix of genes within the chosen pathway in the 2nd set of samples

**Value**

Transcriptional similarity coefficient

**Examples**

```
Pathway1_ExpMat <- matrix(runif(100,0,10), ncol = 10)  
Pathway2_ExpMat <- matrix(runif(100,0,10), ncol = 10)  
TSC(Pathway1_ExpMat, Pathway2_ExpMat)
```

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