## Package 'divergence'

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**Title** Divergence: Functionality for assessing omics data by divergence with respect to a baseline

### Version 1.23.0

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**Description** This package provides functionality for performing divergence analysis as presented in Dinalankara et al, ``Digitizing omics profiles by divergence from a baseline", PANS 2018. This allows the user to simplify high dimensional omics data into a binary or ternary format which encapsulates how the data is divergent from a specified baseline group with the same univariate or multivariate features.

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15

## Contents

| breastTCGA_ER                    | 2  |
|----------------------------------|----|
| breastTCGA_Group                 | 3  |
| breastTCGA_Mat                   | 3  |
| computeChiSquaredTest            | 4  |
| computeMultivariateBinaryMatrix  | 5  |
| computeMultivariateDigitization  | 5  |
| computeMultivariateSupport       | 7  |
| computeQuantileMatrix            | 8  |
| computeUnivariateDigitization    | 9  |
| computeUnivariateSupport         | 10 |
| computeUnivariateTernaryMatrix   | 11 |
| findMultivariateGammaWithSupport | 12 |
| findUnivariateGammaWithSupport   | 13 |
| msigdb_Hallmarks                 | 14 |
|                                  |    |

## Index

breastTCGA\_ER

ER positive or negative status of breast tumor samples

## Description

A factor indicating whether 887 breast samples in breastTCGA\_Mat are ER positive or ER negative. The matched normals have empty values.

## Usage

breastTCGA\_ER

## Format

A Factor of length 887 of levels Negative and Positive (with 111 missing values for the normals).

## Source

https://cancergenome.nih.gov/

breastTCGA\_Group Normal or Tumor status of breast samples

## Description

A factor indicating whether 887 breast samples in breastTCGA\_Mat are tumor or matched normal.

#### Usage

breastTCGA\_Group

## Format

A Factor of length 887 of levels NORMAL and TUMOR.

## Source

https://cancergenome.nih.gov/

breastTCGA\_Mat Gene expression for 260 genes in 887 breast samples

## Description

A data matrix containing a subset of the TCGA breast cancer dataset, with the gene level expression estimates in log2 transcripts per million for 887 breast samples.

## Usage

breastTCGA\_Mat

## Format

A data matrix with 260 rows and 887 columns.

#### Source

https://cancergenome.nih.gov/

computeChiSquaredTest Compute chi-squared test

#### Description

Given a binary or ternary data matrix with class associations of samples, computes chi-squared tests for each feature between given groups

## Usage

```
computeChiSquaredTest(Mat, Groups, classes)
```

#### Arguments

| Mat     | Matrix of digitized binary or ternary data with each column corresponding to a sample and each row corresponding to a feature |
|---------|---|
| Groups  | Factor indicating class association of samples  |
| classes | Vector of class labels; the test will be applied between the classes given.   |

## Value

A data frame with columns 'statistic' and 'pval'.

## Examples

computeMultivariateBinaryMatrix

Compute the binary matrix with digitized divergence coding

#### Description

Function for obtaining the binary form for a matrix for multivariate divergence of data given a baseline range

#### Usage

computeMultivariateBinaryMatrix(seMat, Baseline)

#### Arguments

| seMat    | SummarizedExperiment with assay to be digitized, in [0, 1], with each column corresponding to a sample and each row corresponding to a feature; usually in quantile form. |
|----------|---|
| Baseline | A Baseline object; this corresponds to the output of findMultivariateGammaW-ithSupport() or computeMultivariateSupport()  |

#### Value

A matrix with the same columns as Mat, with rows being the multivariate features, containing the binary form data.

### Examples

```
baseMat = breastTCGA_Mat[, breastTCGA_Group == "NORMAL"]
seMat.base = SummarizedExperiment(assays=list(data=baseMat))
assays(seMat.base)$quantile = computeQuantileMatrix(seMat.base)
baseline = computeMultivariateSupport(seMat=seMat.base, FeatureSets=msigdb_Hallmarks)
dataMat = breastTCGA_Mat[, breastTCGA_Group != "NORMAL"]
seMat = SummarizedExperiment(assays=list(data=dataMat))
assays(seMat)$quantile = computeQuantileMatrix(seMat)
Mat.div = computeMultivariateBinaryMatrix(seMat=seMat, Baseline=baseline)
```

computeMultivariateDigitization *Perform binary digitization* 

#### Description

Function for obtaining the digitized form, along with other relevant statistics and measures given a data matrix and a baseline matrix with multivariate features of interest

#### Usage

```
computeMultivariateDigitization(seMat, seMat.base, FeatureSets,
  computeQuantiles = TRUE, gamma = c(1:9/100, 1:9/10), beta = 0.95,
  alpha = 0.01, distance = "euclidean", verbose = TRUE,
  findGamma = TRUE, Groups = NULL, classes = NULL)
```

## Arguments

| seMat           | SummarizedExperiment with assay to be digitized, in [0, 1], with each column corresponding to a sample and each row corresponding to a feature; usually in quantile form.  |
|-----------------|--|
| seMat.base      | SummarizedExperiment with baseline assay in [0, 1], with each column corresponding to a sample and each row corresponding to a feature   |
| FeatureSets     | The multivariate features in list or matrix form. In list form, each list element<br>should be a vector of individual features; in matrix form, it should be a binary<br>matrix with rownames being individual features and column names being the<br>names of the feature sets. |
| computeQuantile | S  |
|                 | Apply quantile transformation to both data and baseline matrices (TRUE or FALSE; defaults to TRUE).  |
| gamma           | Range of gamma values to search through. By default gamma = $0.01, 0.02,$<br>0.09, 0.1, 0.2,, 0.9.   |
| beta            | Parameter for eliminating outliers ( $0 < beta <= 1$ ). By default beta=0.95.  |
| alpha           | Expected proportion of divergent features per sample to be estimated. The optimal gamma providing this level of divergence in the baseline data will be searched for.  |
| distance        | Type of distance to be calculated between points. Any type of distance that can be passed on to the dist function can be used (default 'euclidean').   |
| verbose         | Logical indicating whether to print status related messages during computation (defaults to TRUE).   |
| findGamma       | Logical indicating whether to search for optimal gamma values through the given gamma values (defaults to TRUE). If FALSE, the first value given in gamma will be used.  |
| Groups          | Factor indicating class association of samples   |
| classes         | Vector of class labels   |

#### Value

A list with elements: Mat.div: divergence coding of data matrix in binary form, of same dimensions at seMat baseMat.div: divergence coding of base matrix in binary form, of same column names at seMat.base, rows being multivariate features. div: data frame with the number of divergent features in each sample features.div: data frame with the divergent probability of each feature; divergence probability for each phenotype in included as well if 'Groups' and 'classes' inputs were provided. Baseline: a list containing a "Ranges" data frame with the baseline interval for each feature, and a "Support" binary matrix of the same dimensions as Mat indicating whether each sample was a support or a feature or not (1=support, 0=not in the support), gamma: selected gamma value alpha: the expected number of divergent features per sample computed over the baseline data matrix

## computeMultivariateSupport

## Examples

```
baseMat = breastTCGA_Mat[, breastTCGA_Group == "NORMAL"]
dataMat = breastTCGA_Mat[, breastTCGA_Group != "NORMAL"]
seMat.base = SummarizedExperiment(assays=list(data=baseMat))
seMat = SummarizedExperiment(assays=list(data=dataMat))
div = computeMultivariateDigitization(
    seMat = seMat,
    seMat.base = seMat.base,
    FeatureSets = msigdb_Hallmarks
)
```

computeMultivariateSupport

Estimate the baseline support

## Description

Function for computing the basline support for multivariate features given gamma and beta parameters.

#### Usage

computeMultivariateSupport(seMat, FeatureSets, gamma = 0.1, beta = 0.95, distance = "euclidean", verbose = TRUE)

## Arguments

| seMat       | SummarizedExperiment with an assay in [0, 1], with each column correspond-<br>ing to a sample and each row corresponding to a feature; usually in quantile<br>form.  |
|-------------|--|
| FeatureSets | The multivariate features in list or matrix form. In list form, each list element<br>should be a vector of individual features; in matrix form, it should be a binary<br>matrix with rownames being individual features and column names being the<br>names of the feature sets. |
| gamma       | Parameter for selecting radius around each support point ( $0 < \text{gamma} < 1$ ). By default gamma = 0.1.   |
| beta        | Parameter for eliminating outliers ( $0 < beta <= 1$ ). By default beta=0.95.  |
| distance    | Type of distance to be calculated between points. Any type of distance that can be passed on to the dist function can be used (default 'euclidean').   |
| verbose     | Logical indicating whether to print status related messages during computation (defaults to TRUE).   |

A list with elements: Support: a matrix indicating which samples were included in the support. Baseline\_list: a list where each element is the baseline of a multivariate feature. featureMat: the multivariate features in matrix form. alpha: the expected number of divergent multivariate features per sample. gamma: the gamma parameter used for baseline computation. distance: the type of distance used for baseline computation.

## Examples

```
baseMat = breastTCGA_Mat[, breastTCGA_Group == "NORMAL"]
seMat.base = SummarizedExperiment(assays=list(data=baseMat))
assays(seMat.base)$quantile = computeQuantileMatrix(seMat.base)
baseline = computeMultivariateSupport(seMat=seMat.base, FeatureSets=msigdb_Hallmarks)
```

computeQuantileMatrix Compute quantile transformations

#### Description

Function for computing the quantile transformation for one or more samples supplied as columns of a matrix.

#### Usage

```
computeQuantileMatrix(seMat)
```

#### Arguments

seMat A data matrix in SummarizedExperiment form, with each column corresponding to a sample and each row corresponding to a feature.

#### Value

A matrix of the same dimensions with the quantile data.

#### Examples

```
baseMat = breastTCGA_Mat[, breastTCGA_Group == "NORMAL"]
seMat.base = SummarizedExperiment(assays=list(data=baseMat))
assays(seMat.base)$quantile = computeQuantileMatrix(seMat.base)
```

8

computeUnivariateDigitization

Perform ternary digitization

## Description

Function for obtaining the digitized form, along with other relevant statistics and measures given a data matrix and a baseline matrix

#### Usage

```
computeUnivariateDigitization(seMat, seMat.base, computeQuantiles = TRUE,
gamma = c(1:9/100, 1:9/10), beta = 0.95, alpha = 0.01,
parallel = TRUE, verbose = TRUE, findGamma = TRUE, Groups = NULL,
classes = NULL)
```

## Arguments

| seMat           | SummarizedExperiment with assay to be digitized, in [0, 1], with each column corresponding to a sample and each row corresponding to a feature; usually in quantile form.       |
|-----------------|---|
| seMat.base      | SummarizedExperiment with baseline assay in [0, 1], with each column corresponding to a sample and each row corresponding to a feature  |
| computeQuantile | es  |
|                 | Logical; apply quantile transformation to both data and baseline matrices (TRUE or FALSE; defaults to TRUE).  |
| gamma           | Range of gamma values to search through. By default gamma = $0.01, 0.02,$ $0.09, 0.1, 0.2,, 0.9$ .  |
| beta            | Parameter for eliminating outliers ( $0 < beta <= 1$ ). By default beta=0.95.   |
| alpha           | Expected proportion of divergent features per sample to be estimated. The optimal gamma providing this level of divergence in the baseline data will be searched for.           |
| parallel        | Logical indicating whether to compute features parallelly with mclapply on<br>Unix based systems (defaults to TRUE, switched to FALSE if parallel package<br>is not available). |
| verbose         | Logical indicating whether to print status related messages during computation (defaults to TRUE).  |
| findGamma       | Logical indicating whether to search for optimal gamma values through the given gamma values (defaults to TRUE). If FALSE, the first value given in gamma will be used.         |
| Groups          | Factor indicating class association of samples (optional).  |
| classes         | Vector of class labels (optional).  |

A list with elements: Mat.div: divergence coding of data matrix in ternary (-1, 0, 1) form, of same dimensions at seMat baseMat.div: divergence coding of base matrix in ternary (-1, 0, 1) form, of same dimensions at seMat.base div: data frame with the number of divergent features in each sample, including upper and lower divergence features.div: data frame with the divergent probability of each feature; divergence probability for each phenotype in included as well if 'Groups' and 'classes' inputs were provided. Baseline: a list containing a "Ranges" data frame with the baseline interval for each feature, and a "Support" binary matrix of the same dimensions as Mat indicating whether each sample was a support or a feature or not (1=support, 0=not in the support), gamma: selected gamma value, alpha: the expected number of divergent features per sample computed over the baseline data matrix, optimal: logical indicaing whether the selected gamma value provided the necessary alpha requirement, alpha\_space: a data frame with alpha values for each gamma searched

#### Examples

```
baseMat = breastTCGA_Mat[, breastTCGA_Group == "NORMAL"]
dataMat = breastTCGA_Mat[, breastTCGA_Group != "NORMAL"]
seMat.base = SummarizedExperiment(assays=list(data=baseMat))
seMat = SummarizedExperiment(assays=list(data=dataMat))
div = computeUnivariateDigitization(
    seMat = seMat,
    seMat.base = seMat.base,
    parallel = TRUE
)
assays(seMat)$div = div$Mat.div
```

computeUnivariateSupport

*Estimate the baseline support* 

#### Description

Function for computing the basline support for univariate features given gamma and beta parameters.

#### Usage

```
computeUnivariateSupport(seMat, gamma = 0.1, beta = 0.95,
    parallel = TRUE, verbose = TRUE)
```

#### Arguments

| seMat | SummarizedExperiment with an assay in [0, 1], with each column correspond-<br>ing to a sample and each row corresponding to a feature; usually in quantile<br>form. |
|-------|---|
| gamma | Parameter for selecting radius around each support point ( $0 < \text{gamma} < 1$ ). By default gamma = 0.1.  |

10

## Value

| beta     | Parameter for eliminating outliers ( $0 < beta <= 1$ ). By default beta=0.95.   |
|----------|---|
| parallel | Logical indicating whether to compute features parallelly with mclapply on<br>Unix based systems (defaults to TRUE, switched to FALSE if parallel package<br>is not available). |
| verbose  | Logical indicating whether to print status related messages during computation (defaults to TRUE).  |

A list with elements "Ranges": data frame with the baseline interval for each feature, "Support": binary matrix of the same dimensions as Mat indicating whether each sample was a support for a feature or not (1=support, 0=not in the support), "gamma": gamma value, and "alpha": the expected number of divergent features per sample estimated over the samples.

## Examples

```
baseMat = breastTCGA_Mat[, breastTCGA_Group == "NORMAL"]
seMat.base = SummarizedExperiment(assays=list(data=baseMat))
assays(seMat.base)$quantile = computeQuantileMatrix(seMat.base)
baseline = computeUnivariateSupport(seMat=seMat.base)
```

computeUnivariateTernaryMatrix

Compute the ternary matrix with digitized divergence coding

## Description

Function for obtaining the ternary form for a matrix of data given a baseline range

#### Usage

```
computeUnivariateTernaryMatrix(seMat, Baseline)
```

#### Arguments

| seMat    | SummarizedExperiment with an assay in [0, 1], with each column correspond-<br>ing to a sample and each row corresponding to a feature; usually in quantile<br>form.                        |
|----------|--|
| Baseline | A list with a data frame element "Ranges" containing the baseline range of each features; this corresponds to the output of findUnivariateGammaWithSupport() or computeUnivariateSupport() |

#### Value

A matrix containing the ternary form data.

## Examples

```
baseMat = breastTCGA_Mat[, breastTCGA_Group == "NORMAL"]
seMat.base = SummarizedExperiment(assays=list(data=baseMat))
assays(seMat.base)$quantile = computeQuantileMatrix(seMat.base)
baseline = computeUnivariateSupport(seMat=seMat.base)
dataMat = breastTCGA_Mat[, breastTCGA_Group != "NORMAL"]
seMat = SummarizedExperiment(assays=list(data=dataMat))
assays(seMat)$quantile = computeQuantileMatrix(seMat)
assays(seMat)$quantile = computeQuantileMatrix(seMat)
```

findMultivariateGammaWithSupport

Find optimal gamma and corresponding support for list of feature sets

## Description

Function for searching through a range of gamma values for finding the smallest gamma and support that provides expected proportion of divergent features per sample less than or equal to alpha.

#### Usage

```
findMultivariateGammaWithSupport(seMat, FeatureSets, gamma = 1:9/10,
    beta = 0.95, alpha = 0.01, distance = "euclidean",
    verbose = TRUE)
```

#### Arguments

| seMat       | SummarizedExperiment with an assay in [0, 1], with each column correspond-<br>ing to a sample and each row corresponding to a feature; usually in quantile<br>form.  |
|-------------|--|
| FeatureSets | The multivariate features in list or matrix form. In list form, each list element<br>should be a vector of individual features; in matrix form, it should be a binary<br>matrix with rownames being individual features and column names being the<br>names of the feature sets. |
| gamma       | Range of gamma values to search through. By default gamma = $\{0.01, 0.02, 0.09, 0.1, 0.2,, 0.9\}$ .   |
| beta        | Parameter for eliminating outliers ( $0 < beta <= 1$ ). By default beta=0.95.  |
| alpha       | Expected proportion of divergent features per sample to be estimated over the samples in Mat. By default alpha = $0.01$ ; i.e. search for the smallest gamma that provides 1% or less number of divergent features per sample.   |
| distance    | Type of distance to be calculated between points. Any type of distance that can be passed on to the dist function can be used (default 'euclidean').   |
| verbose     | Logical indicating whether to print status related messages during computation (defaults to TRUE).   |

12

A list with elements: Support: a matrix indicating which samples were included in the support. Baseline: a list where each element is the baseline of a multivariate feature. featureMat: the multivariate features in matrix form. alpha: the expected number of divergent multivariate features per sample. gamma: the gamma parameter selected. distance: the type of distance used for baselien computation. optimal: TRUE or FALSE indicating whether the alpha criteria was met alpha\_space: the alpha values correspinding to the gamma values searched through

## Examples

```
baseMat = breastTCGA_Mat[, breastTCGA_Group == "NORMAL"]
seMat.base = SummarizedExperiment(assays=list(data=baseMat))
assays(seMat.base)$quantile = computeQuantileMatrix(seMat.base)
baseline = findMultivariateGammaWithSupport(seMat=seMat.base, FeatureSets=msigdb_Hallmarks)
```

findUnivariateGammaWithSupport

Search for optimal gamma and associated support

## Description

Function for searching through a range of gamma values for finding the smallest gamma that provides expected proportion of divergent features per sample less than or equal to alpha.

#### Usage

findUnivariateGammaWithSupport(seMat, gamma = c(1:9/100, 1:9/10), beta = 0.95, alpha = 0.01, parallel = TRUE, verbose = TRUE)

## Arguments

| seMat    | SummarizedExperiment with an assay in [0, 1], with each column correspond-<br>ing to a sample and each row corresponding to a feature; usually in quantile<br>form.  |
|----------|--|
| gamma    | Range of gamma values to search through. By default gamma = $\{0.01, 0.02, 0.09, 0.1, 0.2,, 0.9\}$ .   |
| beta     | Parameter for eliminating outliers ( $0 < beta <= 1$ ). By default beta=0.95.  |
| alpha    | Expected proportion of divergent features per sample to be estimated over the samples in Mat. By default alpha = $0.01$ ; i.e. search for the smallest gamma that provides 1% or less number of divergent features per sample. |
| parallel | Logical indicating whether to compute features parallelly with mclapply on<br>Unix based systems (defaults to TRUE, switched to FALSE if parallel package<br>is not available).  |
| verbose  | Logical indicating whether to print status related messages during computation (defaults to TRUE).   |

A list with elements "Ranges": data frame with the baseline interval for each feature, "Support": binary matrix of the same dimensions as Mat indicating whether each sample was a support for a feature or not (1=support, 0=not in the support), "gamma": gamma value, and "alpha": the expected number of divergent features per sample estimated over the samples, "optimal": logical indicating whether the selected gamma value provided the necessary alpha requirement, and "alpha\_space": a data frame with alpha values for each gamma searched.

## Examples

```
baseMat = breastTCGA_Mat[, breastTCGA_Group == "NORMAL"]
seMat.base = SummarizedExperiment(assays=list(data=baseMat))
assays(seMat.base)$quantile = computeQuantileMatrix(seMat.base)
baseline = findUnivariateGammaWithSupport(seMat=seMat.base)
```

msigdb\_Hallmarks Cancer Hallmark gene sets from the MSigDB collection

#### Description

A subset of the cancer hallmarks functional gene sets from the MSigDB collection.

#### Usage

msigdb\_Hallmarks

## Format

A list of length 10, with the hallmark gene set name, each a character vector of gene symbols.

#### Source

https://http://software.broadinstitute.org/gsea/msigdb/

14

# Index

```
* baseline,
    computeMultivariateSupport, 7
    computeUnivariateSupport, 10
* binary
    computeMultivariateBinaryMatrix, 5
* chi-squared
    computeChiSquaredTest,4
* datasets
    breastTCGA_ER, 2
    breastTCGA_Group, 3
    breastTCGA_Mat, 3
    msigdb_Hallmarks, 14
* digitization
    computeMultivariateBinaryMatrix, 5
    computeUnivariateTernaryMatrix, 11
* digitize
    computeMultivariateDigitization, 5
    computeUnivariateDigitization, 9
* gamma
    findMultivariateGammaWithSupport,
        12
    findUnivariateGammaWithSupport, 13
* support
    computeMultivariateSupport, 7
    computeUnivariateSupport, 10
* ternary
    computeMultivariateDigitization, 5
    computeUnivariateDigitization, 9
    computeUnivariateTernaryMatrix, 11
breastTCGA_ER, 2
breastTCGA_Group, 3
breastTCGA_Mat, 3
computeChiSquaredTest, 4
computeMultivariateBinaryMatrix, 5
computeMultivariateDigitization, 5
computeMultivariateSupport, 7
computeQuantileMatrix, 8
computeUnivariateDigitization, 9
```

computeUnivariateSupport, 10
computeUnivariateTernaryMatrix, 11

findMultivariateGammaWithSupport, 12
findUnivariateGammaWithSupport, 13

msigdb\_Hallmarks, 14