Package 'MiDA'

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Maintainer Elena Filatova <filatova@nniiem.ru>

Description Set of functions designed to simplify transcriptome analysis and identification of marker molecules using microarrays data. The package includes a set of functions that allows performing full pipeline of analysis including data normalization, summarisation, binary classification, FDR (False Discovery Rate) multiple comparison and the definition of potential biological markers.

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Author Elena Filatova [aut, cre], Nikolai Sakharnov [dtc], Dmitry Knyazev [dtc], Oleg Utkin [dtc], Blokhina Scientific Research Institute of Epidemiology and Microbiology of Nizhny Novgorod, Russia [fnd]

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IMexpression

Infectious mononucleosis transcriptome

Description

This is compilied numeric matrix of raw microarray intensities data. Each of 100 rows corresponds to a probe (gene, transcript) and each of 89 column correspondes to a specimen (patient). Specimen is total mRNA samples from human peripheral blood leukocytes' taken from healthy children and children with infectious mononucleosis of different ethiology. Specimen features are in IMspecimen data.

Usage

```
data(IMexpression)
```

Format

A matrix with 100 rows and 89 variables

Details

- rownames. Probes IDs. For most transcripts there are several probes.
- colnames. Specimen IDs.

Author(s)

Nikolai A. Sakharnov, Dmitry I. Knyazev, Oleg V. Utkin

IMspecimen

Description

A dataset containing information about specimens from IMexpression data: IDs and diagnoses.

Usage

data(IMspecimen)

Format

A data frame with 89 rows and 2 variables.

Details

- ID. Specimen ID, factor variable with 89 levels.
- diagnosis. Specimen diagnosis: "ebv" (children with acute EBV mononucleosis), "hhv6" (children with acute HHV6 mononucleosis), "norm" (healthy children). Factor variable with 3 levels.

Author(s)

Nikolai A. Sakharnov, Dmitry I. Knyazev, Oleg V. Utkin

MiBiClassGBODT

Binary classification using gradient boosting over desicion trees

Description

This function conducts a binary classification of specimens based on microarray gene (transcript) expression data. Gradient boosting over desicion trees algorithm is used. Several generalized boosted regression models are fitted during cross-validation, for each model measurements of classification quality and feature importance are returned.

Usage

```
MiBiClassGBODT(Matrix, specimens, n.crossval = 5, ntrees = 10000,
shrinkage = 0.1, intdepth = 2, n.terminal = 10, bag.frac = 0.5)
```

Arguments

Matrix	numeric matrix of expression data where each row corresponds to a probe (gene, transcript), and each column correspondes to a specimen (patient).
specimens	factor vector with two levels specifying specimens in the columns of the Matrix.
n.crossval	integer specifying number of cross-validation folds.
ntrees	integer specifying the total number of decision trees (boosting iterations).
shrinkage	numeric specifying the learning rate. Scales the step size in the gradient descent procedure.
intdepth	integer specifying the maximum depth of each tree.
n.terminal	integer specifying the actual minimum number of observations in the terminal nodes of the trees.
bag.frac	the fraction of the training set observations randomly selected to propose the next tree in the expansion.

Details

Matrix must contain specimens from two classification groups only. To sample expression matrix use MiDataSample.

The order of the variables in specimens and the columns of Matrix must be the same. Levels of specimens are two classification groups. To sample specimens use MiSpecimenSample.

Number of cross-validation folders defines number of models to be fitted. For example, if n.crossval=5 then all specimens are divided into 5 folders, each of them is later used for model testing, so 5 models are fitted. See createFolds for details.

While boosting, basis functions are iteratively adding in a greedy fashion so that each additional basis function further reduces the selected loss function. Gaussian distribution (squared error) is used. ntrees, shrinkage, intdepth are parameters for model tuning. bag.frac introduces randomnesses into the model fit. If bag.frac < 1 then running the same model twice will result in similar but different fits. Number of specimens in train sample must be enough to provide the minimum number of observations in terminal nodes.I.e.

(1-1/n.crossval)*bag.frac > n.terminal.

See gbm for details.

Value

list of 2:

QC - matrix containing quality measures for each fitted model and their summary. Accur - accuracy (percentage of correct predictions), AUC - area under ROC curve (see roc), MCC - Mattew's correlation coefficient

formula ((TP*TN)-(FP*FN))/sqrt((TP+FP)*(TP+FN)*(TN+FP)*(TN+FN)),

F1sc - F1 score

formula 2xPresxRec/(Pres+Rec).

If all the data points from one class are misclassified into other, MCC and F1 score may get NaN values.

Importance - list of data frames containing for each fitted model: var - probe ID and rel.inf - its feature importance for classification (relative influence).

Feature importance (relative influence) graphs are also produced.

MiDataSample

Author(s)

Elena N. Filatova

See Also

createFolds, gbm, MiSpecimenSample, MiDataSample, roc

Examples

MiDataSample	Select matrix columns based	on values of attendant vector
		<i>.</i>

Description

This function selects columns of a matrix that correspond to 1 or 2 factor levels of attendant vector.

Usage

```
MiDataSample(Matrix, specimens, group1, group2 = NULL)
```

Arguments

Matrix	matrix
specimens	factor vector with length equal to number of Matrix columns
group1	value of factor level to sample
group2	additional value of factor level to sample

Details

This function is ment for sampling specimens in gene/transcript expression matrix for binary classification in case when specimens belong to more than two groups. The aim is to create gene/transcript expression matrix that contains specimens for only 1 or 2 groups. Groups are specified in corresponding factor vector that contains specimens description. It should be used together with MiSpecimenSample that samples specimens' description in the same way.

Value

Matrix with rescricted number of columns that correspond to specimens from 1 or 2 groups.

Author(s)

Elena N. Filatova

See Also

MiSpecimenSample

Examples

```
#get gene expression and specimen data
data("IMexpression");data("IMspecimen")
dim(IMexpression) # 100 columns (genes/transcripts) - 89 specimens
colnames(IMexpression)[1:10] # look at first 10 columns of matrix - specimens IDs
IMspecimen[1:10,] # specimens IDs and group factor - diagnoses in attendant vector
# note that specimens in matrix columns are in the same order as specimens in description data
# select specimens with only EBV and NORM diagnoses (and sample the description data as well)
SampleMatrix<-MiDataSample(IMexpression, IMspecimen$diagnosis, "ebv", "norm")
SampleSamples<-MiSpecimenSample(IMspecimen$diagnosis, "ebv", "norm")
dim(SampleMatrix)# only 68 specimens with EBV and NORM diagnoses left
colnames(SampleMatrix)[1:10]
SampleSamples[1:10] # corresponding diagnoses
```

```
MiFracData
```

Subset an expression matrix based on probe's feature importance

Description

This function reduces the number of rows (probes) in gene/transcript expression matrix, leaving only those that have the biggest feature importance for binary classification.

Usage

```
MiFracData(Matrix, importance.list, NRows)
```

Arguments

Matrix	numeric matrix of expression data where each row corresponds to a probe (gene, transcript), and each column correspondes to a specimen (patient). Probe IDs
	must be indicated as matrix row names.
importance.list	
	a list of data frames, containing the result of binary classification: probe IDs in first column and probe's feature importance (relative influence) in the second column in the order from most important to the least important for classification. Such list is the MiBiClassGBODT output (Importance).
NRows	integer defines how many probes are to be left in the expression matrix.

MiFracData

Details

Function provides gene expression matrix subsetting according to probe's feature importance for binary classification, i.e., feature selection. Feature selection provides better classification and identification of significant genes while "not important" genes are taken away from analysis. The procedure of the pairwise combinations of the feature selection and classification methods are described by Pirooznia et al (2008).

The function is able to use multiple feature importance data at a time to subset one expression matrix. If importance.list contains more than one data frame (i.e., the result of a binary classification for more than one model created during cross-validation), the function selects most important probes from each data frame and then removes the repeats. Thus, the output matrix may contain number of probes more than NRows.

Value

expression matrix with only selected probes in alphabetical order as rows and all specimens as columns.

Author(s)

Elena N. Filatova

References

Pirooznia M., Yang J.Y., Yang M.Q., Deng Y. (2008) A comparative study of different machine learning methods on microarray gene expression data. BMC Genomics 9 (Suppl1), S13. https://doi.org/10.1186/1471-2164-9-S1-S13

See Also

MiBiClassGBODT

MiInflCount

Description

Counts mean of probes' feature importance for multiple models of binary classification built on microarray gene/transcript expression data

Usage

```
MiInflCount(importance.list)
```

Arguments

importance.list

a list of data frames, containing the result of binary classification: probe IDs in first column and probe's feature importance (relative influence) in the second column in the order from most important to the least important for classification. Such list is the MiBiClassGBODT output (Importance).

Details

This function takes the result of binary classification performed with cross-validation and counts mean of each probe's feature importance (relative influence) gained in all fitted models.

Value

a list of 2

data.mean - data frame of probe names (in alphabetical order), their mean feature importance and standard deviation.

data.importance - data frame of probe IDs (in alphabetical order) and their original feature importance values gained in all cross-validation models.

Author(s)

Elena N. Filatova

See Also

MiBiClassGBODT

```
# get gene expression and specimen data
data("IMexpression");data("IMspecimen")
# sample expression matrix and specimen data for binary classification,
# only "NORM" and "EBV" specimens are left
SampleMatrix<-MiDataSample(IMexpression, IMspecimen$diagnosis,"norm", "ebv")</pre>
```

MiIntDepthAjust	Ajust maximum depth parameter for fitting generalized boosted regres-
	sion models

Description

Test maximum depth parameter for microarray data binary classification using gradient boosting over desicion trees.

Usage

```
MiIntDepthAjust(Matrix, specimens, test.frac = 5, times = 5,
    ntrees = 1000, shrinkage = 0.1, intdepth = c(1:4),
    n.terminal = 10, bag.frac = 0.5)
```

Arguments

Matrix	numeric matrix of expression data where each row corresponds to a probe (gene, transcript), and each column correspondes to a specimen (patient).
specimens	factor vector with two levels specifying specimens in the columns of the ${\tt Matrix}$
test.frac	integer specifying fraction of data to use for model testing
times	integer specifying number of trials
ntrees	integer specifying the total number of decision trees (boosting iterations).
shrinkage	numeric specifying the learning rate. Scales the step size in the gradient descent procedure.
intdepth	vector of integers specifying the maximum depth of each tree. The tested parameter.
n.terminal	integer specifying the actual minimum number of observations in the terminal nodes of the trees.
bag.frac	the fraction of the training set observations randomly selected to propose the next tree in the expansion.

Details

test.frac defines fraction of specimens that will be used for model testing. For example, if test.frac=5 then 4/5th of specimens will be used for model fitting (train data) and 1/5th of specimens will be used for model testing (test data). Specimens for test and train data will be selected by random. So with times>1, train and test data will differ each time.

While boosting basis functions are iteratively adding in a greedy fashion so that each additional basis function further reduces the selected loss function. Gaussian distribution (squared error) is used. ntrees, shrinkage, intdeep are parameters for model tuning. bag.frac introduces randomnesses into the model fit. If bag.frac < 1 then running the same model twice will result in similar but different fits. Number of specimens in train sample must be enough to provide the minimum number of observations in terminal nodes.I.e.

(1-1/test.frac)*bag.frac > n.terminal.

See gbm for details.

Use MiNTreesAjust and MiShrinkAjust for ajusting other parameters.

Function is rather time-costing. If specimens are not equally distributed between two classified groups, NA may be produced.

Value

list of 2

train.accuracy - a data frame of train data classification accuracy for each intdepth value in each trial and their median.

test.accuracy - a data frame of test data classification accuracy for each intdepth value in each trial and their median.

Also a plot for intdepth versus Accuracy is produced.

Author(s)

Elena N. Filatova

See Also

gbm, MiNTreesAjust, MiShrinkAjust

Examples

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MiNorm

Description

Normalizes microarray expression intensities using different methods with or without background correction.

Usage

MiNorm(Matrix, posNC, method = "none", leaveNC = TRUE, BGcor = FALSE)

Arguments

Matrix	numeric matrix of intensities data where each row corresponds to a probe (gene, transcript), and each column correspondes to a specimen (patient).
posNC	numeric vector specifying numbers of rows containing negative controls (non- coding areas). Used for method="SQN" only. Rows with negative controls will be removed from an intensity matrix after the normalization if leaveNC=FALSE.
method	character string specifying normalization method. Possible values are: "none" (no normalization) "center" (subtracting the row mean), "scale" (dividing by row standard deviation), "standardize" (subtracting the row mean and divid- ing by row standard deviation - z-score transformation), "range" (ranges from 0 to 1), "QN" (normalization based upon quantiles), "SQN" (subset quantile nor- malization using negative control features), "Loess" (cyclicly applying loess normalization).
leaveNC	logical value indicating whether rows with negative control should be deleted from intensity matrix after normalization.
BGcor	logical value indicating whether background correction should be done before normalization. Could be used for background correction only (without data normalization) if method="none".

Details

This function is intended to normalize microarray intensities data between arrays. Background correction is optional.

Background correction method is "normexp", which is based on a convolution model (Ritchie, 2007). See backgroundCorrect for details.

Quantile normalization method implies that we can give each array the same distribution See normalize.quantiles for details.

Subset quantile normalization is performed based on a subset of negative (or non-coding) controls according to (Wu and Aryee, 2010). Number of normal distributions in the mixture approximation is 5, weight given to the parametric normal mixture model is 0.9. See SQN for details.

Cyclic loess normalization implements method of Ballman et al (2004), whereby each array is normalized to the average of all the arrays. See normalizeCyclicLoess for details.

Value

A matrix of the same dimensions as Matrix containing normalized values with or without background correction. If leaveNC=FALSE the function returns a matrix with normalized values without rows containing negative controls.

Author(s)

Elena N. Filatova

References

Ballman K.V., Grill D.E., Oberg A.L. and Therneau T.M. (2004). Faster cyclic loess: normalizing RNA arrays via linear models. Bioinformatics 20, 2778-2786. https://doi.org/10.1093/ bioinformatics/bth327

Bolstad B.M., Irizarry R.A., Astrand M. and Speed T.P. (2003) A Comparison of Normalization Methods for High Density Oligonucleotide Array Data Based on Bias and Variance. Bioinformatics 19(2), 185-193. https://doi.org/10.1093/bioinformatics/19.2.185

Ritchie M.E., Silver J., Oshlack A., Silver J., Holmes M., Diyagama D., Holloway A. and Smyth G.K. (2007). A comparison of background correction methods for two-colour microarrays. Bioinformatics 23, 2700-2707. https://doi.org/10.1093/bioinformatics/btm412

Wu Z and Aryee M. (2010). Subset Quantile Normalization using Negative Control Features. Journal of Computational Biology 17(10), 1385-1395. https://doi.org/10.1089/cmb.2010.0049

See Also

backgroundCorrect, normalizeCyclicLoess, normalize.quantiles, SQN

Examples

```
data("IMexpression")
# Loess normalization
LoMatrix<-MiNorm(IMexpression, method="Loess")
par(mfrow=c(1,2))
boxplot(log2(IMexpression),main="Before normalization")
boxplot(log2(LoMatrix),main="Loess normalization")
par(mfrow=c(1,1))</pre>
```

MiNTreesAjust

Ajust number of trees parameter for fitting generalized boosted regression models

MiNTreesAjust

Description

Test total number of trees parameter for microarray data binary classification using gradient boosting over desicion trees.

Usage

```
MiNTreesAjust(Matrix, specimens, test.frac = 5, times = 5,
    ntrees = c(100, 500, 1000, 5000, 10000), shrinkage = 0.1,
    intdepth = 2, n.terminal = 10, bag.frac = 0.5)
```

Arguments

Matrix	numeric matrix of expression data where each row corresponds to a probe (gene, transcript), and each column correspondes to a specimen (patient).
specimens	factor vector with two levels specifying specimens in the columns of the ${\tt Matrix}$
test.frac	integer specifying fraction of data to use for model testing
times	integer specifying number of trials
ntrees	vector of integer specifying the total number of decision trees (boosting itera- tions).The tested parameter.
shrinkage	numeric specifying the learning rate. Scales the step size in the gradient descent procedure.
intdepth	integer specifying the maximum depth of each tree.
n.terminal	integer specifying the actual minimum number of observations in the terminal nodes of the trees.
bag.frac	the fraction of the training set observations randomly selected to propose the next tree in the expansion.

Details

test.frac defines fraction of specimens that will be used for model testing. For example, if test.frac=5 then 4/5th of specimens will be used for model fitting (train data) and 1/5th of specimens will be used for model testing (test data). Specimens for test and train data will be selected by random. So with times>1, train and test data will differ each time.

While boosting basis functions are iteratively adding in a greedy fashion so that each additional basis function further reduces the selected loss function. Gaussian distribution (squared error) is used. ntrees, shrinkage, intdeep are parameters for model tuning. bag.frac introduces randomnesses into the model fit. If bag.frac < 1 then running the same model twice will result in similar but different fits. Number of specimens in train sample must be enough to provide the minimum number of observations in terminal nodes.I.e.

(1-1/test.frac)*bag.frac > n.terminal.

See gbm for details.

Use MiIntDepthAjust and MiShrinkAjust for ajusting other parameters.

Function is rather time-costing. If specimens are not equally distributed between two classified groups, NA may be produced.

Value

list of 2

train.accuracy - a data frame of train data classification accuracy for each ntrees value in each trial and their median.

test.accuracy - a data frame of test data classification accuracy for each ntrees value in each trial and their median.

Also a plot for ntrees versus Accuracy is produced.

Author(s)

Elena N. Filatova

See Also

gbm, MiIntDepthAjust, MiShrinkAjust

Examples

MiSelectSignif

Select biological markers with high fold change and classification importance

Description

Choose probes which change is biologically significant based on binary classification feature importance, gene expression fold change and statistical significance.

Usage

```
MiSelectSignif(probes, mean1, mean2, FC.method, infl, stat.val,
    tresh.FC = 0.75, tresh.infl = 0.75, tresh.stat = 0.05)
```

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MiSelectSignif

Arguments

probes	character vector of probe (gene, transcript) names.
mean1	numeric vector of mean values for probes expression in the first group of com- parison.
mean2	numeric vector of mean values for probes expression in the second group of comparison.
FC.method	character specifying the method of fold change counting. Possible values are: "absolute" (mean1-mean2), "percent" ((mean1*100/mean2)-100), "ratio" (mean1/mean2), "Log2.ratio" (log2(mean1/mean2)).
infl	numeric vector of mean values for probes feature importance (relative influence) from binary classification.
stat.val	numeric vector of statistical significance (p-value, q-value) for testing differ- ences of mean1 and mean2.
tresh.FC	numeric from o to 1 specifying the tresh hold for fold change FC parameters (quantile). The significant fold change is bigger than tresh. FC.
tresh.infl	numeric from o to 1 specifying the tresh hold for feature importance infl pa- rameters (quantile). The significant feature importance is bigger than tresh.infl.
tresh.stat	numeric from o to 1 specifying the tresh hold for statistical significance stat.val.The significant fold change is lesser than tresh.stat.

Details

The order must be the same for all parameters.

This function marks as "markers" probes that statistically significant change their expression in two groups of comparison with high (over tresh hold) fold change and feature importance from binary classification.

Value

data frame of probe names, their fold change values, statistical significance values, feature inportance values and marker values.

Author(s)

Elena N. Filatova

MiShrinkAjust

Description

Test learning rate (shrinkage) parameter for microarrat data binary classification using gradient boosting over desicion trees.

Usage

```
MiShrinkAjust(Matrix, specimens, test.frac = 5, times = 5,
ntrees = 1000, shrinkage = c(0.001, 0.01, 0.1), intdepth = 2,
n.terminal = 10, bag.frac = 0.5)
```

Arguments

Matrix	numeric matrix of expression data where each row corresponds to a probe (gene, transcript), and each column correspondes to a specimen (patient).
specimens	factor vector with two levels specifying specimens in the columns of the ${\tt Matrix}$
test.frac	integer specifying fraction of data to use for model testing
times	integer specifying number of trials
ntrees	integer specifying the total number of decision trees (boosting iterations).
shrinkage	numeric vector specifying the learning rate. Scales the step size in the gradient descent procedure. The tested parameter.
intdepth	vinteger specifying the maximum depth of each tree.
n.terminal	integer specifying the actual minimum number of observations in the terminal nodes of the trees.
bag.frac	the fraction of the training set observations randomly selected to propose the next tree in the expansion.

Details

test.frac defines fraction of specimens that will be used for model testing. For example, if test.frac=5 then 4/5th of specimens will be used for model fitting (train data) and 1/5th of specimens will be used for model testing (test data). Specimens for test and train data will be selected by random. So with times>1, train and test data will differ each time.

While boosting basis functions are iteratively adding in a greedy fashion so that each additional basis function further reduces the selected loss function. Gaussian distribution (squared error) is used. ntrees, shrinkage, intdeep are parameters for model tuning. bag.frac introduces randomnesses into the model fit. If bag.frac < 1 then running the same model twice will result in similar but different fits. Number of specimens in train sample must be enough to provide the minimum number of observations in terminal nodes.I.e.

(1-1/test.frac)*bag.frac > n.terminal. See gbm for details.

MiSpecimenSample

Use MiIntDepthAjust and MiNTreesAjust for ajusting other parameters.

Function is rather time-costing. If specimens are not equally distributed between two classified groups, NA may be produced.

Value

list of 2

train.accuracy - a data frame of train data classification accuracy for each shrinkage value in each trial and their median.

test.accuracy - a data frame of test data classification accuracy for each shrinkage value in each trial and their median.

Also a plot for shrinkage versus Accuracy is produced.

Author(s)

Elena N. Filatova

See Also

gbm, MiIntDepthAjust, MiNTreesAjust

Examples

MiSpecimenSample Select values from factor vector

Description

This function takes factor vector with multiple levels and selects values for 1 or 2 levels only.

Usage

```
MiSpecimenSample(x, group1, group2 = NULL)
```

Arguments

х	factor vector
group1	value of factor level to sample
group2	additional value of factor level to sample

Details

This function is ment for sampling specimens for binary classification when they belong to more than two groups. The aim is to create factor vector with only two levels specifying specimens in the columns of corresponding microarray expression matrix. It should be used together with MiDataSample that samples columns of microarray expression matrix in the same way.

Value

factor vector with values for 1 or 2 levels only

Author(s)

Elena N. Filatova

See Also

MiDataSample

Examples

```
#get gene expression and specimen data
data("IMexpression");data("IMspecimen")
dim(IMexpression) # 100 columns (genes/transcripts) - 89 specimens
colnames(IMexpression)[1:10] # look at first 10 columns of matrix - specimens IDs
IMspecimen[1:10,] # specimens IDs and group factor - diagnoses in attendant vector
# note that specimens in matrix columns are in the same order as specimens in description data
# select specimens with only EBV and NORM diagnoses (and sample the description data as well)
SampleMatrix<-MiDataSample(IMexpression, IMspecimen$diagnosis, "ebv", "norm")
SampleSamples<-MiSpecimenSample(IMspecimen$diagnosis, "ebv", "norm")
dim(SampleMatrix)# only 68 specimens with EBV and NORM diagnoses left
colnames(SampleMatrix)[1:10]
SampleSamples[1:10] # corresponding diagnoses
```

MiStatCount

FDR for microarray gene expression data

Description

Performs descriptive statistics and FDR (False Discovery Rate) test for microarray expression matrix

MiStatCount

Usage

MiStatCount(Matrix, specimens)

Arguments

Matrix	numeric matrix of expression data where each row corresponds to a probe (gene, transcript), and each column correspondes to a specimen (patient).
specimens	factor vector with two levels specifying specimens in the columns of the Matrix.

Details

This function takes matrix of expression data and performs T-test with FDR correction for two groups for each probe.

T-test is a two-sided, two-class with equal variances against the null hypothesis 'mean1=mean2' for each row. See rowttests for details.

Value

a data frame containing for each probe: mean and sd values for both groups, difference of means, p-value for T-test and q-value for FDR (False Discovery Rate) correction.

Author(s)

Elena N. Filatova

References

Welch B.L.(1951) On the comparison of several mean values: an alternative approach. Biometrika 38, 330-336. https://doi.org/10.1093/biomet/38.3-4.330

See Also

rowttests

```
data("IMexpression"); data("IMspecimen") # load data and specimen information
#sampling data and specimen information
ExpData<-MiDataSample(IMexpression, IMspecimen$diagnosis,"ebv", "norm")
Specimens<-MiSpecimenSample(IMspecimen$diagnosis, "ebv", "norm")
#Counting statistics
StatRes<-MiStatCount(ExpData, Specimens)
head(StatRes)</pre>
```

MiSummarize

Description

Counts median of intensities for multiple probes that target one gene/transcript.

Usage

MiSummarize(Matrix, sep, method = "median")

Arguments

Matrix	numeric matrix of intensities data where each row corresponds to a probe (gene, transcript), and each column correspondes to a specimen (patient). Row names of Matrix should contain probe IDs that consist of three terms: gene name -
	transcript name - probe name.
sep	a character string to separate the terms in probe IDs.
method	character string specifying summarization method. Possible values are "median" and "mean".

Details

This function is used for summarizing expression intensities data when multiple probes target one gene/transcript. Row names of Matrix should contain probe IDs that consist of 3 terms: "Gene name - sep - transcript name - sep - probe name" (for example, "AGTR2.ALL" - for gene, only one probe; "AGTR2.NM_000686.z1" - 1st probe to AGTR2 NM_000686 mRNA transcript).

Value

gene/transcript expression matrix with median/mean of expression intensities for each gene/transcript.

Author(s)

Elena N. Filatova

```
data("IMexpression") # load data
# See 5 zonds to AGTR2.NM_000686
IMexpression [1:10, 1:5]
SumMatrix<-MiSummarize(IMexpression, sep=".")
# now there is median expression for AGTR2.NM_000686
SumMatrix[ 1:10, 1:5]
```

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