

Package ‘biosigner’

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Description Feature selection is critical in omics data analysis to extract restricted and meaningful molecular signatures from complex and high-dimension data, and to build robust classifiers. This package implements a new method to assess the relevance of the variables for the prediction performances of the classifier. The approach can be run in parallel with the PLS-DA, Random Forest, and SVM binary classifiers. The signatures and the corresponding 'restricted' models are returned, enabling future predictions on new datasets. A Galaxy implementation of the package is available within the Workflow4metabolomics.org online infrastructure for computational metabolomics.

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biosigner-package	<i>Molecular signature discovery from omics data</i>
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Description

Feature selection is critical in omics data analysis to extract restricted and meaningful molecular signatures from complex and high-dimension data, and to build robust classifiers. This package implements a new method to assess the relevance of the variables for the prediction performances of the classifier. The approach can be run in parallel with the PLS-DA, Random Forest, and SVM binary classifiers. The signatures and the corresponding 'restricted' models are returned, enabling future predictions on new datasets. A Galaxy implementation of the package is available within the Workflow4metabolomics.org online infrastructure for computational metabolomics.

Author(s)

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See Also

Useful links:

- <http://dx.doi.org/10.3389/fmolb.2016.00026>

biosign*Builds the molecular signature.*

Description

Main function of the 'biosigner' package. For each of the available classifiers (PLS-DA, Random Forest, and SVM), the significant features are selected and the corresponding models are built.

Usage

```
biosign(  
  x,  
  y,  
  methodVc = c("all", "plsda", "randomforest", "svm")[1],  
  bootI = 50,  
  pvalN = 0.05,  
  permI = 1,  
  fixRankL = FALSE,  
  seedI = 123,  
  plotSubC = NA,  
  fig.pdfC = c("none", "interactive", "myfile.pdf")[2],  
  info.txtC = c("none", "interactive", "myfile.txt")[2]  
)
```

S4 method for signature 'matrix'

```
biosign(  
  x,  
  y,  
  methodVc = c("all", "plsda", "randomforest", "svm")[1],  
  bootI = 50,  
  pvalN = 0.05,  
  permI = 1,  
  fixRankL = FALSE,  
  seedI = 123,  
  plotSubC = "",  
  fig.pdfC = c("none", "interactive", "myfile.pdf")[2],  
  info.txtC = c("none", "interactive", "myfile.txt")[2]  
)
```

S4 method for signature 'data.frame'

```
biosign(  
  x,  
  y,  
  methodVc = c("all", "plsda", "randomforest", "svm")[1],  
  bootI = 50,  
  pvalN = 0.05,  
  permI = 1,  
  fixRankL = FALSE,  
  seedI = 123,  
  plotSubC = "",  
  fig.pdfC = c("none", "interactive", "myfile.pdf")[2],
```

```
    info.txtC = c("none", "interactive", "myfile.txt")[2]
  )

## S4 method for signature 'SummarizedExperiment'
biosign(
  x,
  y,
  methodVc = c("all", "plsda", "randomforest", "svm")[1],
  bootI = 50,
  pvalN = 0.05,
  permI = 1,
  fixRankL = FALSE,
  seedI = 123,
  plotSubC = "",
  fig.pdfC = c("none", "interactive", "myfile.pdf")[2],
  info.txtC = c("none", "interactive", "myfile.txt")[2]
)

## S4 method for signature 'MultiAssayExperiment'
biosign(
  x,
  y,
  methodVc = c("all", "plsda", "randomforest", "svm")[1],
  bootI = 50,
  pvalN = 0.05,
  permI = 1,
  fixRankL = FALSE,
  seedI = 123,
  plotSubC = "",
  fig.pdfC = c("none", "interactive", "myfile.pdf")[2],
  info.txtC = c("none", "interactive", "myfile.txt")[2]
)

## S4 method for signature 'ExpressionSet'
biosign(
  x,
  y,
  methodVc = c("all", "plsda", "randomforest", "svm")[1],
  bootI = 50,
  pvalN = 0.05,
  permI = 1,
  fixRankL = FALSE,
  seedI = 123,
  plotSubC = "",
  fig.pdfC = c("none", "interactive", "myfile.pdf")[2],
  info.txtC = c("none", "interactive", "myfile.txt")[2]
)

## S4 method for signature 'MultiDataSet'
biosign(
  x,
  y,
```

```

methodVc = c("all", "plsda", "randomforest", "svm")[1],
bootI = 50,
pvalN = 0.05,
permI = 1,
fixRankL = FALSE,
seedI = 123,
plotSubC = "",
fig.pdfC = c("none", "interactive", "myfile.pdf")[2],
info.txtC = c("none", "interactive", "myfile.txt")[2]
)

```

Arguments

x	Numerical data frame or matrix (observations x variables), or SummarizedExperiment (or ExpressionSet) object ; NAs are allowed for PLS-DA but for SVM, samples with NA will be removed
y	Two-level factor corresponding to the class labels, or a character indicating the name of the column of the pData to be used, when x is an ExpressionSet object
methodVc	Character vector: Either one or all of the following classifiers: Partial Least Squares Discriminant Analysis ('plsda'), or Random Forest ('randomforest'), or Support Vector Machine ('svm')
bootI	Integer: Number of bootstraps for resampling
pvalN	Numeric: To speed up the selection, only variables which significantly improve the model up to two times this threshold (to take into account potential fluctuations) are computed
permI	Integer: Random permutation are used to assess the significance of each new variable included into the model (forward selection)
fixRankL	Logical: Should the initial ranking be computed with the full model only, or as the median of the ranks from the models built on the sampled dataset?
seedI	integer: optional seed to obtain exactly the same signature when rerunning biosigner; default is '123'; set to NULL to prevent seed setting
plotSubC	Character: Graphic subtitle
fig.pdfC	Character: File name with '.pdf' extension for the figure; if 'interactive' (default), figures will be displayed interactively; if 'none', no figure will be generated
info.txtC	Character: File name with '.txt' extension for the printed results (call to sink()); if 'interactive' (default), messages will be printed on the screen; if 'none', no verbose will be generated

Value

An S4 object of class 'biosign' containing the following slots: 1) 'methodVc' character vector: selected classifier(s) ('plsda', 'randomforest', and/or 'svm'), 2) 'accuracyMN' numeric matrix: balanced accuracies for the full models, and the models restricted to the 'S' and 'AS' signatures (predictions are obtained by using the resampling scheme selected with the 'bootI' and 'crossvall' arguments), 3) 'tierMC' character matrix: contains the tier ('S', 'A', 'B', 'C', 'D', or 'E') of each feature for each classifier (features with tier 'S' have been found significant in all backward selections; features with tier 'A' have been found significant in all but the last selection, and so on), 4) modelLs list: selected classifier(s) trained on the subset restricted to the 'S' features, 5) signatureLs list: 'S' signatures for each classifier; and 6) 'AS' list: 'AS' signatures and corresponding trained classifiers, in addition to the dataset restricted to tiers 'S' and 'A' ('xMN') and the labels ('yFc')


```
# restricting to the first 100 features to speed up the example

diaSet <- diaSet[1:100, ]

diaSign <- biosign(diaSet, "type", bootI = 5)
diaSet <- getEset(diaSign)
head(Biobase::fData(diaSet))

detach(diaplasma)

## Application to a MultiAssayExperiment

data("NCI60", package = "ropls")
nci.mae <- NCI60[["mae"]]
library(MultiAssayExperiment)

# Cancer types

table(nci.mae$cancer)

# Restricting to the 'ME' and 'LE' cancer types and to the 'agilent' and 'hgu95' datasets

nci.mae <- nci.mae[, nci.mae$cancer %in% c("ME", "LE"), c("agilent", "hgu95")]

# Selecting the significant features for PLS-DA, RF, and SVM classifiers

nci.mae <- biosign(nci.mae, "cancer", bootI = 5)

# Getting the tiers

SummarizedExperiment::rowData(nci.mae[["agilent"]])

# Getting the models

mae_biosign.ls <- getBiosign(nci.mae)

# Name of the models stored in the (metadata of) each SummarizedExperiment object

names(mae_biosign.ls[["agilent"]])

# Visualizing the individual results

for (set.c in names(mae_biosign.ls))
  plot(mae_biosign.ls[[set.c]][["cancer_plsda.forest.svm"]],
       typeC = "tier",
       plotSubC = set.c)

## Application to a MultiDataSet

data("NCI60", package = "ropls")
nci.mds <- NCI60[["mds"]]

# Restricting to the "agilent" and "hgu95" datasets

nci.mds <- nci.mds[, c("agilent", "hgu95")]

# Restricting to the 'ME' and 'LE' cancer types
```

```

library(Biobase)
sample_names.vc <- Biobase::sampleNames(nci.mds[["agilent"]])
cancer_type.vc <- Biobase::pData(nci.mds[["agilent"]])[, "cancer"]
nci.mds <- nci.mds[sample_names.vc[cancer_type.vc %in% c("ME", "LE")], ]

# Selecting the significant features for PLS-DA, RF, and SVM classifiers

nci_cancer.biosign <- biosign(nci.mds, "cancer", bootI = 5)

# Getting back the updated MultiDataSet

nci.mds <- getMset(nci_cancer.biosign)

```

biosign-class	<i>Class "biosign"</i>
---------------	------------------------

Description

The biosigner object class

Slots

methodVc character vector: selected classifier(s) ('plsda', 'randomforest', or 'svm')

accuracyMN numeric matrix: balanced accuracies for the full models, and the models restricted to the 'S' and 'AS' signatures

tierMC character matrix: contains the tier ('S', 'A', 'B', 'C', 'D', or 'E') of each feature for each classifier

yFc factor with two levels: response factor

modelLs list: selected classifier(s) trained on the subset restricted to the 'S' features

signatureLs list: 'S' signatures for each classifier

xSubMN matrix: dataset restricted to the 'S' tier

AS list: 'AS' signatures and corresponding trained classifiers, in addition to the dataset restricted to tiers 'S' and 'A' ('xMN') and the labels ('yFc')

eset ExpressionSet: when 'biosign' has been applied to an ExpressionSet, the instance with additional columns in fData containing the selected features is stored here

Objects from the Class

Objects can be created by calls of the form `new("biosign", ...)` or by calling the `biosign` function

Author(s)

Philippe Rinaudo and Etienne Thevenot (CEA)

See Also

[biosign](#)

Examples

```
## loading the diaplasm dataset

data(diaplasm)
attach(diaplasm)

## restricting to a smaller dataset for this example

featureSelV1 <- variableMetadata[, "mzmed"] >= 490 & variableMetadata[, "mzmed"] < 500
dataMatrix <- dataMatrix[, featureSelV1]
variableMetadata <- variableMetadata[featureSelV1, ]

## signature selection for all 3 classifiers
## a bootI = 5 number of bootstraps is used for this example
## we recommend to keep the default bootI = 50 value for your analyzes

set.seed(123)
diaSign <- biosign(dataMatrix, sampleMetadata[, "type"], bootI = 5)

detach(diaplasm)
```

biosignMultiDataSet-class

Class "biosignMultiDataSet"

Description

An S4 class to store the biosign objects generated by the application of the 'biosign' method to a MultiDataSet instance

Slots

biosignLs List: List of instances from the 'biosign' class corresponding to the models built on each ExpressionSet

Objects from the Class

Objects can be created by calls of the form `new("biosignMultiDataSet", ...)` or by applying the biosign function to a MultiDataSet instance

See Also

[biosign](#)

Examples

```
# In progress
```

diaplasma*Analysis of plasma from diabetic patients by LC-HRMS*

Description

Plasma samples from 69 diabetic patients were analyzed by reversed-phase liquid chromatography coupled to high-resolution mass spectrometry (Orbitrap Exactive) in the negative ionization mode. The raw data were pre-processed with XCMS and CAMERA (5,501 features), corrected for signal drift, log10 transformed, and annotated with an in-house spectral database. The patient's age, body mass index, and diabetic type are recorded. These three clinical covariates are strongly associated, most of the type II patients being older and with a higher bmi than the type I individuals.

Format

A list with the following elements:

dataMatrix a 69 samples x 5,501 features matrix of numeric type corresponding to the intensity profiles (values have been log10-transformed),

sampleMetadata a 69 x 3 data frame, with the patients' diabetic type ('type', factor), age ('age', numeric), and body mass index ('bmi', numeric),

variableMetadata a 5,501 x 8 data frame, with the median m/z ('mzmed', numeric) and the median retention time in seconds ('rtmed', numeric) from XCMS, the 'isotopes' (character), 'adduct' (character) and 'pcgroups' (numeric) annotations from CAMERA, and the names of the m/z and RT matching compounds from an in-house database of pure spectra from commercial metabolites ('spiDb', character).

Value

List containing the 'dataMatrix' matrix (numeric) of data (samples as rows, variables as columns), the 'sampleMetadata' data frame of sample metadata, and the variableMetadata data frame of variable metadata. Row names of 'dataMatrix' and 'sampleMetadata' are identical. Column names of 'dataMatrix' are identical to row names of 'variableMetadata'. For details see the 'Format' section above.

Source

'diaplasma' dataset.

References

Rinaudo P., Boudah S., Junot C. and Thevenot E.A. (2016). biosigner: a new method for the discovery of significant molecular signatures from omics data. *Frontiers in Molecular Biosciences* 3. doi:10.3389/fmolb.2016.00026

getAccuracyMN	<i>Accuracies of the full model and the models restricted to the signatures</i>
---------------	---

Description

Balanced accuracies for the full models, and the models restricted to the 'S' and 'AS' signatures

Usage

```
getAccuracyMN(object)

## S4 method for signature 'biosign'
getAccuracyMN(object)
```

Arguments

object An S4 object of class biosign, created by the biosign function.

Value

A numeric matrix containing the balanced accuracies for the full models, and the models restricted to the 'S' and 'AS' signatures (predictions are obtained by using the resampling scheme selected with the 'bootI' and 'crossvalI' arguments)

Author(s)

Philippe Rinaudo and Etienne Thevenot (CEA)

Examples

```
## loading the diaplasm dataset

data(diaplasm)
attach(diaplasm)

## restricting to a smaller dataset for this example

featureSelV1 <- variableMetadata[, "mzmed"] >= 490 & variableMetadata[, "mzmed"] < 500
dataMatrix <- dataMatrix[, featureSelV1]
variableMetadata <- variableMetadata[featureSelV1, ]

## signature selection for all 3 classifiers
## a bootI = 5 number of bootstraps is used for this example
## we recommend to keep the default bootI = 50 value for your analyzes

set.seed(123)
diaSign <- biosign(dataMatrix, sampleMetadata[, "type"], bootI = 5)

## individual boxplot of the selected signatures

getAccuracyMN(diaSign)

detach(diaplasm)
```

getBiosign	<i>Getting the biosigner signature from the SummarizedExperiment object</i>
------------	---

Description

The models are extracted as a list

Usage

```
getBiosign(object)

## S4 method for signature 'SummarizedExperiment'
getBiosign(object)

## S4 method for signature 'MultiAssayExperiment'
getBiosign(object)
```

Arguments

object	An S4 object of class SummarizedExperiment, once processed by the biosign method
--------	--

Value

List of biosigner outputs contained in the SummarizedExperiment object

Author(s)

Etienne Thevenot, <etienne.thevenot@cea.fr>

Examples

```
# Getting the diapiasma data set as a SummarizedExperiment

data(diapiasma)

diapiasma.se <- SummarizedExperiment::SummarizedExperiment(assays = list(diapiasma = t(diapiasma[["dataMatrix"]]),
  colData = diapiasma[["sampleMetadata"]]),
  rowData = diapiasma[["variableMetadata"]])

diapiasma.se <- diapiasma.se[1:100, ]

# Selecting the features

diapiasma.se <- biosign(diapiasma.se, "type", bootI = 5, fig.pdfC = "none")

# Getting the signatures

diapiasma.biosign <- getBiosign(diapiasma.se)[["type_plsda_forest_svm"]]

diapiasma.biosign
```

getEset, biosign-method
getEset method

Description

Extracts the complemented ExpressionSet when biosign has been applied to an ExpressionSet

Usage

```
## S4 method for signature 'biosign'  
getEset(object)
```

Arguments

object An S4 object of class biosign, created by biosign function.

Value

An S4 object of class ExpressionSet which contains the dataMatrix (t(exprs(eset))), and the sampleMetadata (pData(eset)) and variableMetadata (fData(eset)) with the additional columns containing the computed tiers for each feature and each classifier.

Author(s)

Etienne Thevenot, <etienne.thevenot@cea.fr>

Examples

```
## loading the diaplasm dataset  
  
data(diaplasm)  
attach(diaplasm)  
  
## building the ExpressionSet instance  
  
diaSet <- Biobase::ExpressionSet(assayData = t(dataMatrix),  
                                phenoData = new("AnnotatedDataFrame",  
                                                  data = sampleMetadata),  
                                featureData = new("AnnotatedDataFrame",  
                                                  data = variableMetadata),  
                                experimentData = new("MIAME",  
                                                     title = "diaplasm"))  
  
## restricting to a smaller dataset for this example  
  
featureSelV1 <- variableMetadata[, "mzmed"] >= 490 & variableMetadata[, "mzmed"] < 500  
diaSet <- diaSet[featureSelV1, ]  
  
## signature selection for all 3 classifiers  
## a bootI = 5 number of bootstraps is used for this example  
## we recommend to keep the default bootI = 50 value for your analyzes
```

```
set.seed(123)
diaSign <- biosign(diaSet, "type", bootI = 5)

diaSet <- biosigner::getEset(diaSign)
head(Biobase::pData(diaSet))
head(Biobase::fData(diaSet))

detach(diaplasma)
```

getMset, biosignMultiDataSet-method
getMset method

Description

Extracts the complemented MultiDataSet when biosign has been applied to a MultiDataSet

Usage

```
## S4 method for signature 'biosignMultiDataSet'
getMset(object)
```

Arguments

object	An S4 object of class <code>biosignMultiDataSet</code> , created by <code>biosign</code> function applied to a <code>MultiDataSet</code>
--------	--

Value

An S4 object of class `MultiDataSet`.

Examples

[illegible]

```

featureData = new("AnnotatedDataFrame",
                  data = fdataDF),
experimentData = new("MIAME",
                    title = setC))

# Adding to the MultiDataSet
nciMset <- MultiDataSet::add_eset(nciMset, eset, dataset.type = setC,
                                GRanges = NA, warnings = FALSE)
}

# Restricting to the 'ME' and 'LE' cancer types
sampleNamesVc <- Biobase::sampleNames(nciMset[["agilent"]])
cancerTypeVc <- Biobase::pData(nciMset[["agilent"]])[, "cancer"]
nciMset <- nciMset[sampleNamesVc[cancerTypeVc %in% c("ME", "LE")], ]
# Summary of the MultiDataSet
nciMset
# Selecting the significant features for PLS-DA, RF, and SVM classifiers, and getting back the updated MultiDataSet
nciBiosign <- biosign(nciMset, "cancer")
nciMset <- getMset(nciBiosign)
# In the updated MultiDataSet, the updated featureData now contains the cancer_biosign_'classifier' columns
# indicating the selected features
lapply(Biobase::fData(nciMset), head)

```

getSignatureLs

*Signatures selected by the models***Description**

List of 'S' (or 'S' and 'A') signatures for each classifier

Usage

```

getSignatureLs(object, tierC = c("S", "AS")[1])

## S4 method for signature 'biosign'
getSignatureLs(object, tierC = c("S", "AS")[1])

```

Arguments

object	An S4 object of class biosign, created by the biosign function.
tierC	Character: defines whether signatures from the 'S' tier only (default) or the ('S' and 'A') tiers should be returned

Value

List of 'S' (or 'S' and 'A') signatures for each classifier

Author(s)

Philippe Rinaudo and Etienne Thevenot (CEA)

Examples

```
## loading the diaplasm dataset

data(diaplasm)
attach(diaplasm)

## restricting to a smaller dataset for this example

featureSelV1 <- variableMetadata[, "mzmed"] >= 490 & variableMetadata[, "mzmed"] < 500
dataMatrix <- dataMatrix[, featureSelV1]
variableMetadata <- variableMetadata[featureSelV1, ]

## signature selection for all 3 classifiers
## a bootI = 5 number of bootstraps is used for this example
## we recommend to keep the default bootI = 50 value for your analyzes

set.seed(123)
diaSign <- biosign(dataMatrix, sampleMetadata[, "type"], bootI = 5)

## individual boxplot of the selected signatures

getSignatureLs(diaSign)

detach(diaplasm)
```

plot,biosign,ANY-method

Plot method for 'biosign' signature objects

Description

Displays classifier tiers or individual boxplots from selected features
 This function plots signatures obtained by biosign.

Usage

```
## S4 method for signature 'biosign,ANY'
plot(
  x,
  y,
  tierMaxC = "S",
  typeC = c("tier", "boxplot")[1],
  plotSubC = "",
  fig.pdfC = c("none", "interactive", "myfile.pdf")[2],
  info.txtC = c("none", "interactive", "myfile.txt")[2]
)

## S4 method for signature 'biosignMultiDataSet,ANY'
plot(
  x,
  y,
```

```

    tierMaxC = "S",
    typeC = c("tier", "boxplot")[1],
    plotSubC = "",
    fig.pdfC = c("none", "interactive", "myfile.pdf")[2],
    info.txtC = c("none", "interactive", "myfile.txt")[2]
  )

```

Arguments

x	An S4 object of class biosign, created by the biosign function.
y	Currently not used.
tierMaxC	Character: Maximum level of tiers to display: Either 'S' and 'A', (for boxplot), or also 'B', 'C', 'D', and 'E' (for tiers) by decreasing number of selections
typeC	Character: Plot type; either 'tier' [default] displaying the comparison of signatures up to the selected 'tierMaxC' or 'boxplot' showing the individual boxplots of the features selected by all the classifiers
plotSubC	Character: Graphic subtitle
fig.pdfC	Character: File name with '.pdf' extension for the figure; if 'interactive' (default), figures will be displayed interactively; if 'none', no figure will be generated
info.txtC	Character: File name with '.txt' extension for the printed results (call to sink()); if 'interactive' (default), messages will be printed on the screen; if 'none', no verbose will be generated

Value

A plot is created on the current graphics device.

Author(s)

Philippe Rinaudo and Etienne Thevenot (CEA)

Examples

```

## loading the diaplasm dataset

data(diaplasm)
attach(diaplasm)

## restricting to a smaller dataset for this example

featureSelV1 <- variableMetadata[, "mzmed"] >= 490 & variableMetadata[, "mzmed"] < 500
dataMatrix <- dataMatrix[, featureSelV1]
variableMetadata <- variableMetadata[featureSelV1, ]

## signature selection for all 3 classifiers
## a bootI = 5 number of bootstraps is used for this example
## we recommend to keep the default bootI = 50 value for your analyzes

diaSign <- biosign(dataMatrix, sampleMetadata[, "type"], bootI = 5)

## individual boxplot of the selected signatures

```

```

plot(diaSign, typeC = "boxplot")

detach(diaplasma)

data("NCI60", package = "ropls")
nci.mds <- NCI60[["mds"]]

# Restricting to the "agilent" and "hgu95" datasets

nci.mds <- nci.mds[, c("agilent", "hgu95")]

# Restricting to the 'ME' and 'LE' cancer types

library(Biobase)
sample_names.vc <- Biobase::sampleNames(nci.mds[["agilent"]])
cancer_type.vc <- Biobase::pData(nci.mds[["agilent"]])[, "cancer"]
nci.mds <- nci.mds[sample_names.vc[cancer_type.vc %in% c("ME", "LE")], ]

# Selecting the significant features for PLS-DA, RF, and SVM classifiers

nci_cancer.biosign <- biosign(nci.mds, "cancer", bootI = 5)
# Plotting the selected signatures
plot(nci_cancer.biosign)

```

predict, biosign-method

Predict method for 'biosign' signature objects

Description

This function predicts values based upon biosign classifiers trained on the 'S' signature

Usage

```
## S4 method for signature 'biosign'
predict(object, newdata, tierMaxC = "S")
```

Arguments

object	An S4 object of class biosign, created by biosign function.
newdata	Either a data frame or a matrix, containing numeric columns only, with column names identical to the 'x' used for model training with 'biosign'.
tierMaxC	Character: Maximum level of tiers to display: Either 'S' or 'A'.

Value

Data frame with the predictions for each classifier as factor columns.

Author(s)

Philippe Rinaudo and Etienne Thevenot (CEA)

Examples

```
## loading the diaplasm dataset

data(diaplasm)
attach(diaplasm)

## restricting to a smaller dataset for this example

featureSelV1 <- variableMetadata[, "mzmed"] >= 490 & variableMetadata[, "mzmed"] < 500
dataMatrix <- dataMatrix[, featureSelV1]
variableMetadata <- variableMetadata[featureSelV1, ]

## training the classifiers
## a bootI = 5 number of bootstraps is used for this example
## we recommend to keep the default bootI = 50 value for your analyzes

set.seed(123)
diaSign <- biosign(dataMatrix, sampleMetadata[, "type"], bootI = 5)

## fitted values (for the subsets restricted to the 'S' signatures)
sFitDF <- predict(diaSign)

## confusion tables
print(lapply(sFitDF, function(predFc) table(actual = sampleMetadata[,
"type"], predicted = predFc)))

## balanced accuracies
sapply(sFitDF, function(predFc) { conf <- table(sampleMetadata[,
"type"], predFc)
conf <- sweep(conf, 1, rowSums(conf), "/")
mean(diag(conf))
})
## note that these values are slightly different from the accuracies
## returned by biosign because the latter are computed by using the
## resampling scheme selected by the bootI or crossvalI arguments
getAccuracyMN(diaSign)["S", ]

detach(diaplasm)
```

show,biosign-method *Show method for 'biosign' signature objects*

Description

Prints the selected features and the accuracies of the classifiers.

Usage

```
## S4 method for signature 'biosign'
show(object)
```

Arguments

object An S4 object of class biosign, created by the biosign function.

Value

Invisible.

Author(s)

Philippe Rinaudo and Etienne Thevenot (CEA)

Examples

```
## loading the diaplasm dataset

data(diaplasm)
attach(diaplasm)

## restricting to a smaller dataset for this example

featureSelV1 <- variableMetadata[, "mzmed"] >= 490 & variableMetadata[, "mzmed"] < 500
dataMatrix <- dataMatrix[, featureSelV1]
variableMetadata <- variableMetadata[featureSelV1, ]

## signature selection for all 3 classifiers
## a bootI = 5 number of bootstraps is used for this example
## we recommend to keep the default bootI = 50 value for your analyzes

set.seed(123)
diaSign <- biosign(dataMatrix, sampleMetadata[, "type"], bootI = 5)

diaSign

detach(diaplasm)
```

SpikePos

Spike-in metabolomics data for apple extracts (from the BioMark package)

Description

Data from a spike-in experiment for apple extracts. Twenty apple extracts are divided in two groups, one control, and one spike-in group. The control group is measured without any spiking - the spike-in group is spiked with nine chemical compounds in three different combinations of concentrations. The data provide the experimental data of the forty apple extracts in the list 'SpikePos' for positive ionization, and in the separate data.frame 'pos.markers' that contains information about the features of the standards, i.e., the spike-in compounds. The data use CAMERA grouping to automatically determine which features are corresponding to which spike-in compounds. Raw data in CDF format are available from the MetaboLights repository (MTBLS59).

Format

'SpikePos' is a list with the following three elements:

data a 40 samples x 1,632 features matrix of numeric type, describing for each of the forty injections the intensity of the features (columns). Column names consist of a combination of retention time (in seconds) and m/z values, and are sorted on retention time,

classes a factor containing the class labels for the forty injections (control, or group1, 2 or 3),
annotation a 1,632 features x 11 metadata data.frame, containing for each of the features XCMS and CAMERA information, such as mz, rt, number of times a feature is identified in the control or spike-in samples, possible isotope or adduct annotation, and whether or not the feature is identified in the standards (the spike-in data).

In addition, 'pos.markers' is a data frame that contains the information of the standards, i.e. the compounds that are spiked in. These data.frames describe in their rows single features identified with XCMS and CAMERA, using the same settings as the experimental apple data, and have the following columns:

comp The (short) name of the spiked-in compound giving rise to this particular feature,
mz, rt, isotope, adduct Feature information, similar to the information in the 'annotation' fields in 'SpikePos',
feature.nr The number of the corresponding feature in 'SpikePos',
group1, group2, group3 Approximate spiking levels for the three groups. A value of 1.0 corresponds to an increase that is roughly equal to the naturally occurring concentration in apple. Exceptions are trans-resveratrol and cyanidin-3-galactoside, both not naturally occurring. These two compounds have been spiked in at one constant level which gives features of comparable size.

Value

'SpikePos' list and 'pos.markers' data frame. For details see the 'Format' section above.

Author(s)

Pietro Franceschi

Source

'SpikePos' dataset.

References

Franceschi, P. et al. (2012) A benchmark spike-in data set for biomarker identification in metabolomics. Journal of Chemometrics, 26, 16–24. DOI:10.1002/cem.1420.

Examples

```
data(SpikePos)
plot(SpikePos$annotation[,c('rt', 'mz')], xlab = 'Time (s)', ylab = 'm/z', main = 'Positive ionization mode')
points(pos.markers[!is.na(pos.markers$feature.nr), c('rt', 'mz')], pch = 19, col = 2)
```

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