# Package 'iC10'

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Title A Copy Number and Expression-Based Classifier for Breast Tumours

Type Package

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Description  Implementation of the classifier described in the paper Ali HR et al (2014) <doi:10.1186 s13059-014-0431-1="">. It uses copy number and/or expression form breast cancer data, trains a Tibshirani's 'pamr' classifier with the features available and predicts the iC10 group.</doi:10.1186>
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iC10-package A Copy Number and Expression-Based Classifier for Breast Tumours

## **Description**

Implementation of the classifier described in the paper Ali HR et al (2014) <doi:10.1186/s13059-014-0431-1>. It uses copy number and/or expression form breast cancer data, trains a Tibshirani's 'pamr' classifier with the features available and predicts the iC10 group.

#### **Details**

#### The DESCRIPTION file:

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Oscar M Rueda <Oscar.Rueda@mrc-bsu.cam.ac.uk> Maintainer:

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iC10 implements the classifier described in the paper 'Genome-driven integrated classification of breast cancer validated in over 7,500 samples' (Ali HR et al., Genome Biology 2014). It uses copy number and/or expression form breast cancer data, trains a pamr classifier (Tibshirani et al.) with the features available and predicts the iC10 group.

#### Author(s)

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Maintainer: Oscar M Rueda < Oscar.Rueda@mrc-bsu.cam.ac.uk>

#### References

Ali HR et al. Genome-driven integrated classification of breast cancer validated in over 7,500 samples. Genome Biology 2014; 15:431. Curtis et al. The genomic and transcriptomic architecture of 2,000 breast tumours reveals novel subgroups. Nature 2012; 486:346-352. Tibshirani et al. Diagnosis of multiple cancer types by shrunken centroids of gene expression. PNAS 2002; 99(10):6567-6572.

# **Examples**

```
require(iC10TrainingData)
data(train.CN)
data(train.Exp)
features <- matchFeatures(Exp=train.Exp, Exp.by.feat="probe")
features <- normalizeFeatures(features, "scale")
res <- iC10(features)
summary(res)
goodnessOfFit(res, newdata=features)
compare(res, iC10=1:2, newdata=features)
compare(res, iC10=2:4, newdata=features)</pre>
```

compare

Compare results of the iC10 classifier

#### **Description**

This function plots the centroids of the training set versus the average profiles of the new data classified in each group.

# Usage

```
compare(obj, iC10=1:10, newdata, name.test="Test",...)
## S3 method for class 'iC10'
compare(obj, iC10=1:10, newdata, name.test="Test",...)
```

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## Arguments

obj An object of class iC10, a result of a call to iC10()

iC10 Groups to plot

ples classified and contained in x. A result of a call to matchFeatures() or

normalizeFeatures()

name.test Name of the new data set to appear in the text of the plot

... Additional arguments passed to plot()

#### Value

A plot is returned with two plots per groups requested.

#### Author(s)

Oscar M. Rueda

#### References

Ali HR et al. Genome-driven integrated classification of breast cancer validated in over 7,500 samples. Genome Biology 2014; 15:431. Curtis et al. The genomic and transcriptomic architecture of 2,000 breast tumours reveals novel subgroups. Nature 2012; 486:346-352.

## See Also

```
iC10, plot.iC10, matchFeatures, normalizeFeatures
```

## **Examples**

```
require(iC10TrainingData)
data(train.CN)
data(train.Exp)
features <- matchFeatures(Exp=train.Exp, Exp.by.feat="probe")
features <- normalizeFeatures(features, "scale")
res <- iC10(features)
compare(res, 1:3, newdata=features)</pre>
```

getCNfeatures

*Internal function for matching copy number features.* 

# **Description**

This function should not be called directly

## Usage

```
getCNfeatures(CN, Probes, Map, by.feat, ref, Synonyms)
```

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# **Arguments**

CN CN features matrix

Probes Vector with the probes to match

Map data.frame with the genomic description of the features to match

by . feat "probe" or "gene", indicating if match should be done by probe position or gene

name.

ref hg18 or hg19 (only relevant if matching is done by probe position).

Synonyms data.frame with available synonym gene names to match (only relevant if match-

ing is done by gene name).

## Value

A matrix with the copy number features

## Author(s)

Oscar M Rueda

getExpfeatures

Internal function for matching expression features.

# **Description**

Internal function for matching expression features.

# Usage

```
getExpfeatures(Exp, Probes, Synonyms, by.feat)
```

## **Arguments**

Exp Matrix of expression features
Probes Vector of probes to match

Synonyms vector of synonyms fo gene names

by.feat either "probe" or "gene"

## Value

A matrix with the Probes in Exp.

## Note

This function is not supposed to be called directly. use matchFeatures instead.

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## Author(s)

Oscar M Rueda

#### References

Curtis et al. The genomic and transcriptomic architecture of 2,000 breast tumours reveals novel subgroups. Nature 2012; 486:346-352.

#### See Also

matchFeatures

goodnessOfFit

Goodness of fit results of the iC10 classifier

## **Description**

Goodness of fit results of the iC10 classifier: this function computes correlations between the signatures of the training dataset and the classified features.

# Usage

```
goodnessOfFit(obj, iC10=1:10, newdata=NULL,...)
## S3 method for class 'iC10'
goodnessOfFit(obj, iC10=1:10, newdata=NULL,...)
```

#### Arguments

obj An object of iC10 class.

iC10 Groups to compute goodness of fit.

newdata The feature data to compute the goodness of fit. Must be the samples classified

in obj. It can be a call to matchFeatures or normalizeFeatures. If NULL,

obj\$fitted is used.

... Additional arguments passed to cor (like method; Default is pearson)

## Value

It prints the correlation for each iC10.

# Author(s)

Oscar M Rueda

#### References

Ali HR et al. Genome-driven integrated classification of breast cancer validated in over 7,500 samples. Genome Biology 2014; 15:431. Curtis et al. The genomic and transcriptomic architecture of 2,000 breast tumours reveals novel subgroups. Nature 2012; 486:346-352.

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## See Also

iC10

#### **Examples**

```
require(iC10TrainingData)
data(train.CN)
data(train.Exp)
features <- matchFeatures(Exp=train.Exp, Exp.by.feat="probe")
features <- normalizeFeatures(features, "scale")
res <- iC10(features)
goodnessOfFit(res, newdata=features)</pre>
```

iC10

A copy number and expression-based classfier for breast cancers

# **Description**

iC10 implements the classifier described in the paper 'Genome-driven integrated classification of breast cancer validated in over 7,500 samples' (Ali HR et al., Genome Biology 2014). It uses copy number and/or expression form breast cancer data, trains a pamr classifier (Tibshirani et al.) with the features available and predicts the iC10 group.

## Usage

```
iC10(x, seed=25435)
```

## **Arguments**

x An object with class iC10features: A list with elements 'train.CN', 'train.Exp',

'train.iC10', 'CN', 'Exp', 'map.cn', 'map.exp'

seed to initialize random number generator. It is passed to set.seed(). See

details.

#### **Details**

This function trains a pamr classifier and predicts the set of samples. The shrinkage parameter is obtained with crossvalidation, therefore different runs can give different results (unless a seed is specified).

## Value

An object of class iC10. A list with the following elements:

class Prediction classes for the samples

posterior Probablitites for each sample to belong to each of the 10 groups

centroids Shrunken Centroids for each of the 10 groups.

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fitted	Normalized features for the samples classified.
map.cn	Annotation data for the copy number features
map.exp	Annotation data for the expression features

## Author(s)

Oscar M. Rueda

#### References

Ali HR et al. Genome-driven integrated classification of breast cancer validated in over 7,500 samples. Genome Biology 2014; 15:431. Curtis et al. The genomic and transcriptomic architecture of 2,000 breast tumours reveals novel subgroups. Nature 2012; 486:346-352. Tibshirani et al. Diagnosis of multiple cancer types by shrunken centroids of gene expression. PNAS 2002; 99(10):6567-6572.

## See Also

See pamr. train, pamr. cv and pamr. predict in package pamr.

# **Examples**

```
require(iC10TrainingData)
data(train.CN)
data(train.Exp)
features <- matchFeatures(Exp=train.Exp, Exp.by.feat="probe")
features <- normalizeFeatures(features, "scale")
res <- iC10(features)</pre>
```

matchFeatures

Matching features from the classifier to the test data.

# Description

This function matches available copy number and/or expression data features to the training signatures; using either genomic position or HUGO gene name for copy number features and either Illumina probe names or HUGO gene name for expression features.

# Usage

```
matchFeatures(CN = NULL, Exp = NULL,
CN.by.feat = c("gene", "probe"),
Exp.by.feat = c("gene", "probe"),
ref="hg19")
```

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#### **Arguments**

CN Data must be log2 copy number ratios. Two formats are allowed: - a matrix

where each row represents a gene and each column a sample. In this case CN.by.feat must be "gene" and the rownames must be the hgnc gene names. - a data.frame with segmented data. The following columns must exist: 'ID' for the sample name, 'chromosome\_name' for the chromosome (must be numeric), 'loc.start' for the start position of the region, 'loc.end' for the end position of the region, 'seg.mean' for the log2ratio of the segment. If NULL, copy number is

not used in the classifier.

Exp Matrix with the expression data to classify. Each row must be a gene or an

Illumina probe, and each column must correspond to a sample. Rownames must be either Illumina probes, in which case Exp.by.feat must be "probe"; or hgnc gene names, in which case Exp.by.feat must be "gene". If NULL, expression

is not used in the classifier.

CN.by.feat Either "probe" or "gene", Default is "probe".

Exp.by.feat Either "probe" or "gene", Default is "gene".

ref Either "hg18", "hg19" or "hg38". It is used to match the copy number probes if

CN.by.feat is "probe"

## **Details**

One of CN or Exp must be not NULL. If matching is done by gene, hgnc gene name is used to match the rownames of the features. A list of synonym gene names is used (see Map.All). For copy number features matched by probe, the maximum log ratio in absolute value inside the limits of the feature is used. If there is no copy number in that region, the value of the probe before it is used.

# Value

A list with the following elements is returned:

CN copy number data to classify train.CN copy number training data

Exp expression data to classify train.Exp expression training data

train.iC10 iC10 assignments for the training data
map.cn annotation data for the copy number features
map.exp annotation data for the expression features

#### Note

Note that the training set will be different, depending on the features matched. Genomic annotation for the training dataset has been obtained from Mark Dunning's lluminaHumanv3.db package.

## Author(s)

Oscar M Rueda

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## References

Ali HR et al. Genome-driven integrated classification of breast cancer validated in over 7,500 samples. Genome Biology 2014; 15:431. Curtis et al. The genomic and transcriptomic architecture of 2,000 breast tumours reveals novel subgroups. Nature 2012; 486:346-352.

## See Also

```
normalizeFeatures, iC10
```

## **Examples**

```
require(iC10TrainingData)
data(train.CN)
data(train.Exp)
features <- matchFeatures(Exp=train.Exp,Exp.by.feat="probe", ref="hg18")
str(features)</pre>
```

normalizeFeatures

Normalization of expression features

# **Description**

Normalization of expression features. Several methods available in the package CONOR can be used.

## Usage

```
normalizeFeatures(x, method=c("none", "scale"))
```

# **Arguments**

x An object result of a call to matchFeatures

method Several methods are available: "none": No normalization is done "scale": Each

expression feature is scaled to have zero mean and standard deviation 1

#### **Details**

No further normalization is needed on the copy number, as log2 ratios are comparable between platforms.

# Value

A list of the same format as matchFeatures, but with train. Exp anfd Exp normalized.

## Note

As CONOR package is no longer maintained, the methods are not available temporarily. We will include more normalization methods in the next version of this package.

plot.iC10

#### Author(s)

Oscar M Rueda

#### References

Ali HR et al. Genome-driven integrated classification of breast cancer validated in over 7,500 samples. Genome Biology 2014; 15:431. Curtis et al. The genomic and transcriptomic architecture of 2,000 breast tumours reveals novel subgroups. Nature 2012; 486:346-352.

# **Examples**

```
require(iC10TrainingData)
data(train.CN)
data(train.Exp)
features <- matchFeatures(Exp=train.Exp,
Exp.by.feat="probe", ref="hg18")
features <- normalizeFeatures(features, "scale")</pre>
```

plot.iC10

Plot results of the iC10 classifier

## **Description**

Plot results of the iC10 classifier, in two different formats: either the signatures of the training set or the signatures of the new data classified.

# Usage

```
## S3 method for class 'iC10'
plot(x, sample.name=1, newdata = NULL,...)
```

## **Arguments**

x An object of iC10 class:

sample.name Number of sample to plot (if newdata is NULL). It can be either a number or a

character with the sample name.

newdata An object result to call to matchFeatures or normalizeFeatures containing

the features of the samples to plot.

... Additional arguments passed to plot.

#### **Details**

Two types of plots can be produced. If newdata is NULL, a panel 6x2 is drawn with the 10 profiles of the signatures of the training set and the profile of the features of sample. name and the distribution of the probabilities of classification to each iC10 for that sample. If newdata is not nutll, a panel 6x2 (with the 11th panel empty) is drawn with the 10 profiles of newdata samples and their distribution into the clusters. The features are sorted by type: copy number (if available) are drawn in grey, and then expression, each of them are sorted by genomic position.

print.iC10

# Value

A 6x2 plot is produced.

#### Author(s)

Oscar M Rueda

#### References

Ali HR et al. Genome-driven integrated classification of breast cancer validated in over 7,500 samples. Genome Biology 2014; 15:431. Curtis et al. The genomic and transcriptomic architecture of 2,000 breast tumours reveals novel subgroups. Nature 2012; 486:346-352.

## See Also

iC10

## **Examples**

```
require(iC10TrainingData)
data(train.CN)
data(train.Exp)
features <- matchFeatures(Exp=train.Exp, Exp.by.feat="probe")
features <- normalizeFeatures(features, "scale")
res <- iC10(features)
plot(res, sample.name=10)
plot(res, newdata=features)</pre>
```

print.iC10

Print results of the iC10 classifier

# Description

Print results of the iC10 classifier

# Usage

```
## S3 method for class 'iC10'
print(x, ...)
```

# **Arguments**

x An object of iC10 class:

... Additional arguments passed to print.

# Value

It returns a call to str.

summary.iC10

## Author(s)

Oscar M Rueda

#### References

Ali HR et al. Genome-driven integrated classification of breast cancer validated in over 7,500 samples. Genome Biology 2014; 15:431. Curtis et al. The genomic and transcriptomic architecture of 2,000 breast tumours reveals novel subgroups. Nature 2012; 486:346-352.

#### See Also

iC10

#### **Examples**

```
require(iC10TrainingData)
data(train.CN)
data(train.Exp)
features <- matchFeatures(Exp=train.Exp, Exp.by.feat="probe")
features <- normalizeFeatures(features, "scale")
res <- iC10(features)
res</pre>
```

summary.iC10

Summary results of the iC10 classifier

# **Description**

Summary results of the iC10 classifier: shows the distribution of samples classified into each iC10 group and a summary of the maximum posterior probablity for each sample. Small values pinpoint samples with no clear group assigned.

## Usage

```
## S3 method for class 'iC10'
summary(object, ...)
```

# **Arguments**

object An object of iC10 class.
... Additional arguments passed to summary.

## Value

The function prints a table of the classification ad a summary of the maximum posterior probability for each sample.

summary.iC10

## Author(s)

Oscar M Rueda

#### References

Ali HR et al. Genome-driven integrated classification of breast cancer validated in over 7,500 samples. Genome Biology 2014; 15:431. Curtis et al. The genomic and transcriptomic architecture of 2,000 breast tumours reveals novel subgroups. Nature 2012; 486:346-352. Tibshirani et al. Diagnosis of multiple cancer types by shrunken centroids of gene expression. PNAS 2002; 99(10):6567-6572.

# See Also

See iC10 and pamr.train, pamr.cv and pamr.predict in package pamr.

# **Examples**

```
require(iC10TrainingData)
data(train.CN)
data(train.Exp)
features <- matchFeatures(Exp=train.Exp,
Exp.by.feat="probe", ref="hg18")
features <- normalizeFeatures(features, "scale")
res <- iC10(features)
summary(res)</pre>
```

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