

Package ‘roccv’

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Type Package

Title ROC for Cross Validation Results

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Depends R (>= 3.0.0), glmnet, parallel, pROC

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Description Cross validate large genetic data while specifying clinical variables that should always be in the model using the function cv(). An ROC plot from the cross validation data with AUC can be obtained using rocplot(), which also can be used to compare different models. Framework was built to handle genetic data, but works for any data.

ByteCompile TRUE

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NeedsCompilation no

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cv*Cross validation results for a model*

Description

Cross validation results for a model

Usage

```
cv(clinical_x = NULL, genomic_x = NULL, y = NULL, data = NULL,
  clinical_formula = NULL, family = "binomial", folds = NULL, k = 10,
  fit_method = "glm", method_name = NULL, n.cores = 1, ...)
```

Arguments

clinical_x	clinical variables that will always be included in the model
genomic_x	genomic variables that will be penalized if a penalized model is used
y	response variables
data	dataframe if clinical formula is used
clinical_formula	formula for clinical variables
family	gaussian, binomial or poisson
folds	predefined partitions for cross validation
k	number of cross validation folds. A value of k=n is leave one out cross validation.
fit_method	glm or glmnet used to fit the model
method_name	tracking variable to include in return dataframe
n.cores	Number of cores to be used
...	additional commands to glm or cv.glmnet

Value

returns a dataframe of predicted values and observed values. In addition, method_name is recorded if that variable is defined.

Author(s)

Ben Sherwood <ben.sherwood@ku.edu>

Examples

```
x <- matrix(rnorm(800),ncol=8)
y <- runif(100) <- exp(1 + x[,1] + x[,5])/(1+exp(1 + x[,1] + x[,5]))
cv_results <- cv(x,y=y,method_name="without_formula")
combined_data <- data.frame(y=y,x1=x[,1],x5=x[,5])
gx <- x[,c(2,3,4,6,7,8)]
cvf <- cv(genomic_x=gx,clinical_formula=y~x1+x5,data=combined_data,method_name="with_form")
```

fit_pred_fold	<i>Cross validation on fold i</i>
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Description

Cross validation on fold i

Usage

```
fit_pred_fold(i, x, y, folds, fit_method, family, non_pen_vars = NULL, ...)
```

Arguments

i	target partition
x	matrix of predictors
y	vector of responses
folds	defines how data is separated into folds for cross validation
fit_method	model being used to fit the data
family	family used to fit the data
non_pen_vars	index of variables that will not be penalized if glmnet is used
...	additional commands to glm or cv.glmnet

Value

returns predictions for partition i

Author(s)

Ben Sherwood <ben.sherwood@ku.edu>

Examples

```
folds_10 <- randomly_assign(100,10)
x <- matrix(rnorm(800),ncol=8)
y <- runif(100) <- exp(1 + x[,1] + x[,5])/(1+exp(1 + x[,1] + x[,5]))
fold_1_results <- fit_pred_fold(1,x,y,folds_10,"glm","binomial")
fold_2_results <- fit_pred_fold(2,x,y,folds_10,"glm","binomial")
```

randomly_assign	<i>Assigns n samples into k groups</i>
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Description

Assigns n samples into k groups

Usage

```
randomly_assign(n, k)
```

Arguments

n	sample size
k	number of groups

Value

returns a vector of length n with a random assignment of entries from 1 to k

Author(s)

Ben Sherwood <ben.sherwood@ku.edu>

Examples

```
n <- 100
folds_10 <- randomly_assign(n,10)
folds_5 <- randomly_assign(n,5)
```

roccv	<i>roccv: A package for creating ROC plots on cross validated data</i>
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Description

roccv: A package for creating ROC plots on cross validated data

rocplot*Create ROC plot from cross validation results*

Description

Create ROC plot from cross validation results

Usage

```
rocplot(plot_data, ...)
```

Arguments

plot_data	dataframe with columns: response, prediction and method
...	additional commands plot.roc such as main

Value

returns ROC plot

Author(s)

Ben Sherwood <ben.sherwood@ku.edu>

Examples

```
x <- matrix(rnorm(800),ncol=8)
y <- runif(100) < exp(1 + x[,1] + x[,5])/(1+exp(1 + x[,1] + x[,5]))
cv_results <- cv(x,y=y,method_name="without_formula")
combined_data <- data.frame(y=y,x1=x[,1],x5=x[,5])
gx <- x[,c(2,3,4,6,7,8)]
cvf <- cv(genomic_x=gx,clinical_formula=y~x1+x5,
data=combined_data,method_name="with_form")
total_results <- rbind(cv_results,cvf)
rocplot(total_results,main="rocplot test")
```

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