Package ‘sperrorest’

February 20, 2015

Type Package
Title Spatial Error Estimation and Variable Importance
Version 0.2-1
Date 2012-06-19
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Depends ROCR, rpart
Suggests ipred, RSAGA, sp
Description This package implements spatial error estimation and
permutation-based variable importance measures for predictive
models using spatial cross-validation and spatial block
bootstrap.
License GPL-2
LazyLoad yes
Collate 'sperrorest-package.R' 'sperrorest.R' 'sperrorest_misc.R'
'sperrorest_resampling.R' 'sperrorest_error.R'
'sperrorest_data.R'
Repository CRAN
Date/Publication 2012-06-20 07:10:59
NeedsCompilation no

R topics documented:

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Description

This package implements spatial error estimation and permutation-based spatial variable importance using different spatial cross-validation and spatial block bootstrap methods. To cite ‘sperrorest’ in publications, reference the paper by Brenning (2012).

References


### Description

Add distance information to resampling objects

### Usage

```r
add.distance(object, ...)  
```

```r
## S3 method for class 'resampling'
add.distance(object, data,  
coords = c("x", "y"), ...)
```

```r
## S3 method for class 'represampling'
add.distance(object, ...)
```

### Arguments

- `object`: `resampling` or `represampling` object
- `...`: Additional arguments to `dataset.distance` and `add.distance.resampling`, respectively
- `data`: data frame containing at least the columns specified by `coords`
- `coords`: (ignored by `partition.cv`)

### Details

Nearest-neighbour distances are calculated for each sample in the test set. These `nrow(test)` nearest-neighbour distances are then averaged. Aggregation methods other than `mean` can be chosen using the `fun` argument, which will be passed on to `dataset.distance`.

### Value

A `resampling` or `represampling` object containing an additional `$distance` component in each `resampling` object. The distance component is a single numeric value indicating, for each `train` / `test` pair, the (by default, mean) nearest-neighbour distance between the two sets.

### See Also

`dataset.distance`, `represampling`, `resampling`
Examples

```r
data(ecuador) # Muenchow et al. (2012), see ?ecuador
ns.parti = partition.cv(ecuador)
sp.parti = partition.kmeans(ecuador)
ns.parti = add.distance(ns.parti, ecuador)
sp.parti = add.distance(sp.parti, ecuador)
# non-spatial partitioning: very small test-training distance:
ns.parti[[1]][[1]]$distance
# spatial partitioning: more substantial distance, depending on number of folds etc.
sp.parti[[1]][[1]]$distance
```

---

### as.represampling

**Resampling objects with repetition, i.e. sets of partitionings or bootstrap samples**

#### Description

Functions for handling represampling objects, i.e. lists of `resampling` objects.

#### Usage

```r
as.represampling(object, ...) 
```

#### Arguments

- `object` object of class `represampling`, or a list to be coerced to this class
- `x` object of class `resampling`
- `...` currently not used

#### Details

Represampling objects are (names) lists of `resampling` objects. Such objects are typically created by `partition.cv`, `partition.kmeans`, `represampling.disc.bootstrap` and related functions.

In r-repeated k-fold cross-validation, for example, the corresponding represampling object has length r, and each of its r `resampling` objects has length k.

as.resampling.list coerces object to class represampling while coercing its elements to `resampling` objects. Some validity checks are performed.
as.resampling

Value

as.represampling methods return an object of class represampling with the contents of object.

See Also

resampling, partition.cv, partition.kmeans, represampling, disc.bootstrap, etc.

Examples

data(ecuador) # Muenchow et al. (2012), see ?ecuador
# Partitioning by elevation classes in 200 m steps:
fac = factor( as.character( floor( ecuador$deg / 300 ) ) )
summary(fac)
parti = as.resampling(fac)
# a list of lists specifying sets of training and test sets,
# using each factor at a time as the test set:
str(parti)
summary(parti)

---

as.resampling  Resampling objects such as partitionings or bootstrap samples

Description

Create/coerce and print resampling objects, e.g., partitionings or bootstrap samples derived from a data set.

Usage

as.resampling(object, ...)

## Default S3 method:
as.resampling(object, ...)

## S3 method for class 'factor'
as.resampling(object, ...)

## S3 method for class 'list'
as.resampling(object, ...)

validate.resampling(object)
is.resampling(x, ...)

## S3 method for class 'resampling'
print(x, ...)
Arguments

object: depending on the function/method, a list or a vector of type factor defining a partitioning of the dataset

x: object of class resampling

Details

A resampling object is a list of lists defining a set of training and test samples.

In the case of k-fold cross-validation partitioning, for example, the corresponding resampling object would be of length k, i.e. contain k lists. Each of these k lists defines a training set of size \(n(k-1)/k\) (where \(n\) is the overall sample size), and a test set of size \(n/k\). The resampling object does, however, not contain the data itself, but only indices between 1 and \(n\) identifying the selection (see Examples).

Another example is bootstrap resampling. `represampling.bootstrap` with argument `oob=TRUE` generates [rep]resampling objects with indices of a bootstrap sample in the train component and indices of the out-of-bag sample in the test component (see Examples below).

`as.resampling.factor`: For each factor level of the input variable, `as.resampling.factor` determines the indices of samples in this level (= test samples) and outside this level (= training samples). Empty levels of object are dropped without warning.

`as.resampling.list` checks if the list in object has a valid resampling object structure (with components `train` and `test` etc.) and assigns the class attribute "resampling" if successful.

Value

as.resampling methods: An object of class resampling.

See Also

`represampling, partition.cv, partition.kmeans, represampling.bootstrap`, etc.

Examples

```r
data(ecuador) # Muenchow et al. (2012), see ?ecuador

# Partitioning by elevation classes in 200 m steps:
parti = factor( as.character( floor( ecuador$dem / 200 ) ) )
smp = as.resampling(parti)
summary(smp)
# Compare:
summary(parti)

# k-fold (non-spatial) cross-validation partitioning:
parti = partition.cv(ecuador)
parti = parti[[1]] # the first (and only) resampling object in parti
# data corresponding to the test sample of the first fold:
str( ecuador[ parti[[1]]$test , ] )
# the corresponding training sample - larger:
```

```
as.tilename

```
str( ecuador[ parti[[1]]$train , ] )

# Bootstrap training sets, out-of-bag test sets:
pari = represampling.bootstrap(ecuador, oob = TRUE)
pari = pari[[1]] # the first (and only) resampling object in pari
# out-of-bag test sample: approx. one-third of nrow(ecuador):
str( ecuador[ pari[[1]]$test , ] )
# bootstrap training sample: same size as nrow(ecuador):
str( ecuador[ pari[[1]]$train , ] )
```

---

**as.tilename**

*Alphanumeric tile names*

**Description**

Functions for generating and handling alphanumeric tile names of the form "X2:Y7" as used by `partition.tiles` and `represampling.tile.bootstrap`.

**Usage**

```
as.tilename(x, ...)

## S3 method for class 'numeric'
as.tilename(x, ...)

## S3 method for class 'tilename'
as.character(x, ...)

## S3 method for class 'tilename'
as.numeric(x, ...)

## S3 method for class 'character'
as.tilename(x, ...)

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

**Arguments**

- **x**
  - object of class tilename, character, or numeric (of length 2)
- **...**
  - additional arguments (currently ignored)

**Value**

object of class tilename, character, or numeric vector of length 2

**See Also**

`partition.tiles`, `represampling.represampling.tile.bootstrap`
dataset.distance

Example

tnm = as.tilename(c(2,3))
tnm # "X2:Y3"
as.numeric(tnm) # c(2,3)

dataset.distance

Description

dataset.distance calculates Euclidean nearest-neighbour distances between two point datasets and summarizes these distances using some function, by default the mean.

Usage

dataset.distance(d1, d2, x.name = "x", y.name = "y",
fun = mean, method = "euclidean", ...) 

Arguments

d1 a data.frame with (at least) columns with names given by x.name and y.name; these contain the x and y coordinates, respectively

d2 see d1 - second set of points

x.name name of column in d1 and d2 containing the x coordinates of points

y.name same for y coordinates

fun function to be applied to the vector of nearest-neighbor distances of d1 from d2

method type of distance metric to be used; only 'euclidean' is currently supported

Details

Nearest-neighbour distances are calculated for each point in d1, resulting in a vector of length nrow(d1), and fun is applied to this vector.

Value

depends on fun; typically (e.g., mean) a numeric vector of length 1

See Also

add.distance

Examples

d = data.frame(x = rnorm(100), y = rnorm(100))
dataset.distance(d, d) # == 0
**ecuador** *J. Muenchow's Ecuador landslide data set*

**Description**

Data set created by Jannes Muenchow, University of Erlangen-Nuremberg, Germany. These data should be cited as Muenchow et al. (2012) (see reference below). This publication also contains additional information on data collection and the geomorphology of the area. The data set provided here is (a subset of) the one from the 'natural' part of the RBSF area and corresponds to landslide distribution in the year 2000.

**Format**

a data.frame with point samples of landslide and non-landslide locations in a study area in the Andes of southern Ecuador.

**References**


**Examples**

```r
data(ecuador)
str(ecuador)
library(rpart)
ctrl = rpart.control(cp = 0.02)
fit = rpart(slides ~ dem + slope + hcurv + vcurv +
            log.carea + cslope,
            data = ecuador, control = ctrl)
par(xpd = TRUE)
plot(fit, compress = TRUE, main = "Muenchow's landslide data set")
text(fit, use.n = TRUE)
```

---

**err.default** *Default error function*

**Description**

Calculate a variety of accuracy measures from observations and predictions of numerical and categorical response variables.

**Usage**

```r
err.default(obs, pred)
```
get.small.tiles

Arguments

obs  factor, logical, or numeric vector with observations
pred factor, logical, or numeric vector with predictions. Must be of same type as obs with the exception that pred may be numeric if obs is factor or logical ('soft' classification).

Value

A list with (currently) the following components, depending on the type of prediction problem:

'hard' classification
misclassification error, overall accuracy; if two classes, sensitivity, specificity, positive predictive value (PPV), negative predictive value (NPV), kappa

'soft' classification
area under the ROC curve, error and accuracy at a obs>0.5 dichotomization, false-positive rate (FPR; 1-specificity) at 70, 80 and 90 percent sensitivity, true-positive rate (sensitivity) at 80, 90 and 95 percent specificity

regression bias, standard deviation, mean squared error, MAD (mad), median, interquartile range (IQR) of residuals

Note

NA values are currently not handled by this function, i.e. they will result in an error.

See Also

ROCR

Examples

obs = rnorm(1000)
# Two mock (soft) classification examples:
err.default( obs>0, rnorm(1000) ) # just noise
err.default( obs>0, obs + rnorm(1000) ) # some discrimination
# Three mock regression examples:
err.default( obs, rnorm(1000) ) # just noise, but no bias
err.default( obs, obs + rnorm(1000) ) # some association, no bias
err.default( obs, obs + 1 ) # perfect correlation, but with bias

get.small.tiles Identify small partitions that need to be fixed.

Description

get.small.tiles identifies partitions (tiles) that are too small according to some defined criterion / criteria (minimum number of samples in tile and/or minimum fraction of entire dataset).
Usage

get.small.tiles(tile, min.n = NULL, min.frac = 0,
ignore = c())

Arguments

tile factor: tile/partition names for all samples; names must be coercible to class tilename, i.e. of the form "X4:Y2" etc.
min.n integer (optional): minimum number of samples per partition
min.frac numeric >0, <1: minimum relative size of partition as percentage of sample
ignore character vector: names of tiles to be ignored, i.e. to be retained even if the inclusion criteria are not met.

Value

character vector: names of tiles that are considered 'small' according to these criteria

See Also

partition.tiles, tilename

Examples

data(ecuador) # Muenchow et al. (2012), see ?ecuador
# Rectangular partitioning without removal of small tiles:
parti = partition.tiles(ecuador, nsplit = c(10,10), reassign = FALSE)
summary(parti)
length(parti[[1]])
# Same in factor format for the application of get.small.tiles:
parti.fac = partition.tiles(ecuador, nsplit = c(10,10), reassign = FALSE, return.factor = TRUE)
get.small.tiles(parti.fac[[1]], min.n = 20) # tiles with less than 20 samples
par2 = partition.tiles(ecuador, nsplit = c(10,10), reassign = TRUE,
min.n = 20, min.frac = 0)
length(par2[[1]]) < length(parti[[1]])

partition.cv Partition the data for a (non-spatial) cross-validation

Description

partition.cv creates a represampling object for length(repetition)-repeated nfold-fold cross-validation.

Usage

partition.cv(data, coords = c("x", "y"), nfold = 10,
repetition = 1, seed1 = NULL, return.factor = FALSE)
Arguments

- **data**: data.frame containing at least the columns specified by coords
- **coords**: (ignored by partition.cv)
- **nfold**: number of partitions (folds) in nfold-fold cross-validation partitioning
- **repetition**: numeric vector: cross-validation repetitions to be generated. Note that this is not the number of repetitions, but the indices of these repetitions. E.g., use repetition = c(1:100) to obtain (the 'first') 100 repetitions, and repetition = c(101:200) to obtain a different set of 100 repetitions.
- **seed**: seed[i] is the random seed that will be used by set.seed in repetition i (i in repetition) to initialize the random number generator before sampling from the data set.
- **return.factor**: if FALSE (default), return a represampling object; if TRUE (used internally by other sperrorest functions), return a list containing factor vectors (see Value)

Details

This function does not actually perform a cross-validation or partition the data set itself; it simply creates a data structure containing the indices of training and test samples.

Value

If return.factor=FALSE (the default), a represampling object. Specifically, this is a (named) list of length(repetition) resampling objects. Each of these resampling objects is a list of length nfold corresponding to the folds. Each fold is represented by a list of containing the components train and test, specifying the indices of training and test samples (row indices for data). If return.factor=TRUE (mainly used internally), a (named) list of length length(repetition). Each component of this list is a vector of length nrow(data) of type factor, specifying for each sample the fold to which it belongs. The factor levels are factor(1:nfold).

See Also

sperrorest, represampling

Examples

```r
data(ecuador)
## non-spatial cross-validation:
resamp = partition.cv(ecuador, nfold = 5, repetition = 1:2)
plot(resamp, ecuador)
# first repetition, second fold, test set indices:
idx = resamp[["1"]][[2]]$test
# test sample used in this particular repetition and fold:
ecuador[idx,]
```
Description

`partition.cv.strat` creates a set of sample indices corresponding to cross-validation test and training sets.

Usage

```r
partition.cv.strat(data, coords = c("x", "y"),
                   nfold = 10, return.factor = FALSE, repetition = 1,
                   seed = NULL, strat)
```

Arguments

- `coords` vector of length 2 defining the variables in `data` that contain the x and y coordinates of sample locations
- `strat` character: column in `data` containing a factor variable over which the partitioning should be stratified; or factor vector of length `nrow(data)`: variable over which to stratify
- `data` `data.frame` containing at least the columns specified by `coords`
- `nfold` number of partitions (folds) in `nfold`-fold cross-validation partitioning
- `return.factor` if `FALSE` (default), return a `represampling` object; if `TRUE` (used internally by other `sperrorest` functions), return a list containing factor vectors (see Value)
- `repetition` numeric vector: cross-validation repetitions to be generated. Note that this is not the number of repetitions, but the indices of these repetitions. E.g., use `repetition = c(1:100)` to obtain the 'first' 100 repetitions, and `repetition = c(101:200)` to obtain a different set of 100 repetitions.
- `seed` `seed += i` is the random seed that will be used by `set.seed` in repetition `i` (`i` in `repetition`) to initialize the random number generator before sampling from the data set.

Value

A `represampling` object, see also `partition.cv`, `partition.strat.cv`, however, stratified with respect to the variable `data[,strat]`; i.e., cross-validation partitioning is done within each set `data[data[,strat]==i,](i in levels(data[,strat]))`, and the `i`th folds of all levels are combined into one cross-validation fold.

See Also

`sperrorest, as.resampling, resample.strat.uniform`
Examples

data(ecuador)
parti = partition.cv.strat(ecuador, strat = "slides", nfold = 5, repetition = 1)
idx = parti[["1"]][[1]]$train
mean(ecuador$slides[idx]=="TRUE") / mean(ecuador$slides=="TRUE")
# always == 1
# Non-stratified cross-validation:
parti = partition.cv(ecuador, nfold = 5, repetition = 1)
idx = parti[["1"]][[1]]$train
mean(ecuador$slides[idx]=="TRUE") / mean(ecuador$slides=="TRUE")
# close to 1 because of large sample size, but with some random variation

---

partition.disc  Leave-one-disc-out cross-validation and leave-one-out cross-validation

Description

partition.disc partitions the sample into training and tests set by selecting circular test areas (possibly surrounded by an exclusion buffer) and using the remaining samples as training samples (leave-one-disc-out cross-validation). partition.loo creates training and test sets for leave-one-out cross-validation with (optional) buffer.

Usage

partition.disc(data, coords = c("x", "y"), radius, buffer = NULL, ndisc = nrow(data), seed1 = NULL, return.train = TRUE, prob = NULL, replace = FALSE, repetition = 1)

partition.loo(data, ndisc = nrow(data), replace = FALSE, ...)

Arguments

coops  vector of length 2 defining the variables in data that contain the x and y coordinates of sample locations
radius  radius of test area discs; performs leave-one-out resampling if radius <0
buffer  radius of additional 'neutral area' around test area discs that is excluded from training and test sets; defaults to 0, i.e. all samples are either in the test area or in the training area.
ndisc  Number of discs to be randomly selected; each disc constitutes a separate test set. Defaults to nrow(data), i.e. one disc around each sample.
return.train  If FALSE, returns only test sample; if TRUE, also the training area.
prob  optional argument to sample
replace  optional argument to sample: sampling with or without replacement?
partition.factor

repetition see partition.cv; however, see Note below: repetition should normally be =1 in this function.

... arguments to be passed to partition.disc

data data.frame containing at least the columns specified by coords

seed seed1+i is the random seed that will be used by set.seed in repetition i (i in repetition) to initialize the random number generator before sampling from the data set.

Value

A represampling object. Contains length(repetition) resampling objects. Each of these contains n.disc lists with indices of test and (if return.train=TRUE) training sets.

Note

Test area discs are centered at (random) samples, not at general random locations. Test area discs may (and likely will) overlap independently of the value of replace. replace only controls the replacement of the center point of discs when drawing center points from the samples. radius<0 does leave-one-out resampling with an optional buffer. radius=0 is similar except that samples with identical coordinates would fall within the test area disc.

References


See Also

sperrorest, partition.cv, partition.kmeans

Examples

data(ecuador)
parti = partition.disc(ecuador, radius=200, buffer=200, n.disc=5, repetition=1:2)
plot(parti,ecuador)
summary(parti)
# leave-one-out with buffer:
parti.loo = partition.loo(ecuador, buffer=200)
summary(parti)

partition.factor

Partition the data for a (non-spatial) leave-one-factor-out cross-validation based on a given, fixed partitioning

Description

partition.factor creates a represampling object, i.e. a set of sample indices defining cross-validation test and training sets.
Usage

```r
partition.factor(data, coords = c("x", "y"), fac, return.factor = FALSE, repetition = 1)
```

Arguments

- `coords`: vector of length 2 defining the variables in `data` that contain the x and y coordinates of sample locations.
- `fac`: either the name of a variable (column) in `data`, or a vector of type factor and length `nrow(data)` that contains the partitions to be used for defining training and test samples.
- `data`: `data.frame` containing at least the columns specified by `coords`.
- `return.factor`: if `FALSE` (default), return a `represampling` object; if `TRUE` (used internally by other `sperrorest` functions), return a list containing factor vectors (see Value).
- `repetition`: numeric vector: cross-validation repetitions to be generated. Note that this is not the number of repetitions, but the indices of these repetitions. E.g., use `repetition=c(1:100)` to obtain (the 'first') 100 repetitions, and `repetition=c(101:200)` to obtain a different set of 100 repetitions.

Value

A `represampling` object, see also `partition.cv` for details.

Note

In this partitioning approach, all repetitions are identical and therefore pseudo-replications.

See Also

`sperrorest`, `partition.cv`, `as.resampling.factor`

Examples

```r
data(ecuador)
# I don't recommend using this partitioning for cross-validation,
# this is only for demonstration purposes:
breaks = quantile(ecuador$dem, seq(0,1,length=6))
ecuador$zclass = cut(ecuador$dem, breaks, include.lowest=TRUE)
summary(ecuador$zclass)
parti = partition.factor(ecuador, fac = "zclass")
plot(parti,ecuador)
summary(parti)
```
Description

partition.kmeans divides the study area into irregularly shaped spatial partitions based on \textit{k}-means (\texttt{kmeans}) clustering of spatial coordinates.

Usage

\begin{verbatim}
partition.kmeans(data, coords = c("x", "y"), nfold = 10,
repetition = 1, seed1 = NULL, return.factor = FALSE,
balancing.steps = 1, order.clusters = TRUE, ...)
\end{verbatim}

Arguments

- \texttt{coords}: vector of length 2 defining the variables in \texttt{data} that contain the x and y coordinates of sample locations
- \texttt{nfold}: number of cross-validation folds, i.e. parameter \textit{k} in \textit{k}-means clustering
- \texttt{balancing.steps}: if >1, perform \texttt{nfold}-means clustering \texttt{balancing.steps} times, and pick the clustering that minimizes the Gini index of the sample size distribution among the partitions. The idea is that 'degenerate' partitions will be avoided, but this also has the side effect of reducing variation among partitioning repetitions. More meaningful constraints (e.g., minimum number of positive and negative samples within each partition should be added in the future.
- \texttt{order.clusters}: if \texttt{TRUE}, clusters are ordered by increasing x coordinate of center point
- \texttt{...}: additional arguments to \texttt{kmeans}
- \texttt{data}: \texttt{data.frame} containing at least the columns specified by \texttt{coords}
- \texttt{repetition}: numeric vector: cross-validation repetitions to be generated. Note that this is not the number of repetitions, but the indices of these repetitions. E.g., use \texttt{repetition=c(1:100)} to obtain (the 'first') 100 repetitions, and \texttt{repetition=c(101:200)} to obtain a different set of 100 repetitions.
- \texttt{seed1}: \texttt{seed1+i} is the random seed that will be used by \texttt{set.seed} in \texttt{repetition} \texttt{i} (\texttt{i} in \texttt{repetition}) to initialize the random number generator before sampling from the data set.
- \texttt{return.factor}: if \texttt{FALSE} (default), return a \texttt{represampling} object; if \texttt{TRUE} (used internally by other \texttt{sperrorest} functions), return a list containing factor vectors (see Value)

Value

A \texttt{represampling} object, see also \texttt{partition.cv} for details.

Note

Default parameter settings may change in future releases.
partition.tiles

References


See Also

sperrorest, partition.cv, partition.disc, partition.tiles, kmeans

Examples

data(ecuador)
resamp = partition.kmeans(ecuador, nfold = 5, repetition = 1:2)
plot(resamp, ecuador)

Description

partition.tiles divides the study area into a specified number of rectangular tiles. Optionally small partitions can be merged with adjacent tiles to achieve a minimum number or percentage of samples in each tile.

Usage

partition.tiles(data, coords = c("x", "y"),
    dsplit = NULL, nsplit = NULL,
    rotation = c("none", "random", "user"), user.rotation,
    offset = c("none", "random", "user"), user.offset,
    reassign = TRUE, min.frac = 0.025, min.n = 5,
    iterate = 1, return.factor = FALSE, repetition = 1,
    seed1 = NULL)

Arguments

coords vector of length 2 defining the variables in data that contain the x and y coordinates of sample locations

dsplit optional vector of length 2: equidistance of splits in (possibly rotated) x direction (dsplit[1]) and y direction (dsplit[2]) used to define tiles. If dsplit is of length 1, its value is recycled. Either dsplit or nsplit must be specified.

nsplit optional vector of length 2: number of splits in (possibly rotated) x direction (nsplit[1]) and y direction (nsplit[2]) used to define tiles. If nsplit is of length 1, its value is recycled.
rotation indicates whether and how the rectangular grid should be rotated; random rotation is only between -45 and +45 degrees.

user.rotation if rotation="user", angles (in degrees) by which the rectangular grid is to be rotated in each repetition. Either a vector of same length as repetition, or a single number that will be replicated length(repetition) times.

offset indicates whether and how the rectangular grid should be shifted by an offset

user.offset if offset="user", a list (or vector) of two components specifying a shift of the rectangular grid in (possibly rotated) x and y direction. The offset values are relative values, a value of 0.5 resulting in a one-half tile shift towards the left, or upward. If this is a list, its first (second) component refers to the rotated x (y) direction, and both components must have same length as repetition (or length 1). If a vector of length 2 (or list components have length 1), the two values will be interpreted as relative shifts in (rotated) x and y direction, respectively, and will therefore be recycled as needed (length(repetition) times each).

reassign logical (default TRUE): if TRUE, 'small' tiles (as per min.frac and min.n arguments and get.small.tiles) are merged with (smallest) adjacent tiles. If FALSE, small tiles are 'eliminated', i.e. set to NA.

min.frac numeric >=0, <1: minimum relative size of partition as percentage of sample; argument passed to get.small.tiles. Will be ignored if NULL.

min.n integer >=0: minimum number of samples per partition; argument passed to get.small.tiles. Will be ignored if NULL.

iterate argument to be passed to tile.neighbors

data data.frame containing at least the columns specified by coords

return.factor if FALSE (default), return a resampling object; if TRUE (used internally by other sparrowrest functions), return a list containing factor vectors (see Value)

repetition numeric vector: cross-validation repetitions to be generated. Note that this is not the number of repetitions, but the indices of these repetitions. E.g., use repetition=c(1:100) to obtain (the 'first') 100 repetitions, and repetition=c(101:200) to obtain a different set of 100 repetitions.

seed seed+i is the random seed that will be used by set.seed in repetition i (i in repetition) to initialize the random number generator before sampling from the data set.

Value

A resampling object. Contains length(repetition) resampling objects as repetitions. The exact number of folds / test-set tiles within each resampling objects depends on the spatial configuration of the data set and possible cleaning steps.

Note

Default parameter settings may change in future releases. This function, especially the rotation and shifting part of it and the algorithm for cleaning up small tiles is still a bit experimental. Use with caution. For non-zero offsets (offset!="none"), the number of tiles may actually be greater than nsplit[1]*nsplit[2] because of fractional tiles lurking into the study region. reassign=TRUE with suitable thresholds is therefore recommended for non-zero (including random) offsets.
plot.represampling

Description

plot.represampling displays the partitions or samples corresponding arising from the resampling of a data set.

Usage

## S3 method for class 'represampling'
plot(x, data,
    coords = c("x", "y"), pch = "+", wiggle.sd = 0, ...)

## S3 method for class 'resampling'
plot(x, ...)

Arguments

x a represampling resp. resampling object
data a data.frame of samples containing at least the x and y coordinates of samples as specified by coords
coords vector of length 2 defining the variables in data that contain the x and y coordinates of sample locations
pch point symbol (to be passed to points)
wiggle.sd 'wiggle' the point locations in x and y direction to avoid overplotting of samples drawn multiple times by bootstrap methods; this is a standard deviation (in the units of the x/y coordinates) of a normal distribution and defaults to 0 (no wigging)

See Also

sperrorest, as.resampling.factor, get.small.tiles, tile.neighbors

Examples

data(ecuador)
parti = partition.tiles(ecuador, nsplit = c(4,3), reassign = FALSE)
plot(parti, ecuador)
summary(parti) # tile A4 has only 55 samples
# same partitioning, but now merge tiles with less than 100 samples to adjacent tiles:
parti2 = partition.tiles(ecuador, nsplit = c(4,3), reassign = TRUE, min.n = 100)
plot(parti2, ecuador)
summary(parti2)
# tile B4 (in 'parti') was smaller than A3, therefore A4 was merged with B4, not with A3
# now with random rotation and offset, and tiles of 2000 m length:
parti3 = partition.tiles(ecuador, dsplit = 2000, offset = "random", rotation = "random", reassign = TRUE, min.n = 100)
plot(parti3, ecuador)
summary(parti3)
Note

This function is not intended for samples obtained by resampling with replacement (e.g., bootstrap) because training and test points will be overplotted in that case. The size of the plotting region will also limit the number of maps that can be displayed at once, i.e., the number of rows (repetitions) and fields (columns).

Examples

data(ecuador)
# non-spatial cross-validation:
  resamp = partition.cv(ecuador, nfold = 5, repetition = 1:2)
  plot(resamp, ecuador)
# spatial cross-validation using k-means clustering:
  resamp = partition.kmeans(ecuador, nfold = 5, repetition = 1:2)
  plot(resamp, ecuador)

represampling.bootstrap

Non-spatial bootstrap resampling

Description

represampling.bootstrap draws a bootstrap random sample (with replacement) from data.

Usage

represampling.bootstrap(data, coords = c("x", "y"),
nboot = nrow(data), repetition = 1, seed1 = NULL,
oob = FALSE)

Arguments

coops vector of length 2 defining the variables in data that contain the x and y coordinates of sample locations

nboot Size of bootstrap sample

oob logical (default FALSE): if TRUE, use the out-of-bag sample as the test sample; if FALSE, draw a second bootstrap sample of size nboot independently to obtain a test sample

data data.frame containing at least the columns specified by coords

repetition numeric vector: cross-validation repetitions to be generated. Note that this is not the number of repetitions, but the indices of these repetitions. E.g., use repetition=c(1:100) to obtain the 'first' 100 repetitions, and repetition=c(101:200) to obtain a different set of 100 repetitions.

seed1 seed1+i is the random seed that will be used by set.seed in repetition i (i in repetition) to initialize the random number generator before sampling from the data set.
Value

A represampling object. This is a (named) list containing length(repetition) resampling objects. Each of these contains only one list with indices of training and test samples. Indices are row indices for data.

Examples

```r
data(ecuador)
# only 10 bootstrap repetitions, normally use >=100:
parti = represampling.bootstrapping(ecuador, repetition = 10)
plot(parti, ecuador) # careful: overplotting occurs
# because some samples are included in both the training and
# the test sample (possibly even multiple times)
```

Represampling.disc.bootstrap

Overlapping spatial block bootstrap using circular blocks

Description

Represampling.disc.bootstrap performs a spatial block bootstrap by resampling at the level of rectangular partitions or 'tiles' generated by partition.tiles.

Usage

```r
represampling.disc.bootstrap(data, coords = c("x", "y"),
nboot, repetition = 1, seed1 = NULL, oob = FALSE, ...)
```

Arguments

- oob logical (default FALSE): if TRUE, use the out-of-bag sample as the test sample (the complement of the nboot[1] test set discs, minus the buffer area as specified in the ... arguments to partition.disc); if FALSE, draw a second bootstrap sample of size nboot independently to obtain a test sample (sets of overlapping discs drawn with replacement)
- nboot number of bootstrap samples; you may specify different values for the training sample (nboot[1]) and for the test sample (nboot[2])
- ... additional arguments to be passed to partition.disc; note that a buffer argument has not effect if oob=FALSE; see example below
- data data.frame containing at least the columns specified by coords
- coords vector of length 2 defining the variables in data that contain the x and y coordinates of sample locations
- repetition numeric vector: cross-validation repetitions to be generated. Note that this is not the number of repetitions, but the indices of these repetitions. E.g., use repetition=c(1:100) to obtain the 'first' 100 repetitions, and repetition=c(101:200) to obtain a different set of 100 repetitions.
seed\_1 $$+ i$$ is the random seed that will be used by `set.seed` in repetition $$i$$ ($$i$$ in repetition) to initialize the random number generator before sampling from the data set.

**Note**

Performs $$n_{boot}$$ out of $$n_{row(data)}$$ resampling of circular discs. This is an *overlapping* spatial block bootstrap where the blocks are circular.

**Examples**

```r
data(ecuador)
# Overlapping disc bootstrap:
parti = represampling.factor.bootstrap(ecuador, radius=200, nboot=20, oob=FALSE)
plot(parti, ecuador)
# Note that a 'buffer' argument would make no difference because bootstrap sets of discs are
drawn independently for the training and test sample.
#
# Overlapping disc bootstrap for training sample, out-of-bag sample as test sample:
parti = represampling.factor.bootstrap(ecuador, radius=200, buffer=200, nboot=10, oob=TRUE)
plot(parti, ecuador)
```

**Description**

`represampling.factor.bootstrap` resamples partitions defined by a factor variable. This can be used for non-overlapping block bootstraps and similar.

**Usage**

```r
represampling.factor.bootstrap(data, fac, repetition = 1,
nboot = -1, seed1 = NULL, oob = FALSE)
```

**Arguments**

- `fac` defines a grouping or partitioning of the samples in data; three possible types: (1) the name of a variable in data (coerced to factor if not already a factor variable); (2) a factor variable (or a vector that can be coerced to factor); (4) a list of factor variables (or vectors that can be coerced to factor); this list must be of length `length(repetition)`, and if it is named, the names must be equal to `as.character(repetition)`; this list will typically be generated by a partition.*/ function with `return.factor=TRUE` (see Examples below)
represampling.kmeans.bootstrap

nboot number of bootstrap replications used for generating the bootstrap training sample (nboot[1]) and the test sample (nboot[2]); nboot[2] is ignored (with a warning) if oob=TRUE. A value of -1 will be substituted with the number of levels of the factor variable, corresponding to an \( n \) out of \( n \) bootstrap at the grouping level defined by fac.

oob if TRUE, the test sample will be the out-of-bag sample; if FALSE (default), the test sample is an independently drawn bootstrap sample of size nboot[2]

data data.frame containing at least the columns specified by coords

repetition numeric vector: cross-validation repetitions to be generated. Note that this is not the number of repetitions, but the indices of these repetitions. E.g., use repetition=c(1:100) to obtain the first 100 repetitions, and repetition=c(101:200) to obtain a different set of 100 repetitions.

seed seed+i is the random seed that will be used by set.seed in repetition i (i in repetition) to initialize the random number generator before sampling from the data set.

Details

nboot refers to the number of groups (as defined by the factors) to be drawn with replacement from the set of groups. I.e., if fac is a factor variable, nboot would normally not be greater than nlevels(fac), nlevels(fac) being the default as per nboot=-1.

See Also

represampling.disc.bootstrap, represampling.tile.bootstrap, note yet implemented: partition.cv.factor

Examples

data(ecuador)
# a dummy example for demonstration, performing bootstrap
# at the level of an arbitrary factor variable:
parti = represampling.factor.bootstrap(ecuador, factor(floor(ecuador$dem/100)), oob=TRUE)
plot(parti,ecuador)
# using the factor bootstrap for a non-overlapping block bootstrap
# (see also represampling.tile.bootstrap):
fac = partition.tiles(ecuador, return.factor=TRUE, repetition=c(1:3), dsplit=500, min.n=200, rotation="random",
parti = represampling.factor.bootstrap(ecuador, fac, oob=TRUE, repetition=c(1:3))
plot(parti,ecuador)

represampling.kmeans.bootstrap

Spatial block bootstrap at the level of spatial k-means clusters

Description

represampling.kmeans.bootstrap performs a non-overlapping spatial block bootstrap by resampling at the level of irregularly-shaped partitions generated by partition.kmeans.
represampling.tile.bootstrap

Usage

represampling.kmeans.bootstrap(data,
   coords = c("x", "y"), repetition = 1, nfold = 10,
   nboot = nfold, seed1 = NULL, oob = FALSE, ...)

Arguments

nfold see partition.kmeans
nboot see represampling.factor.bootstrap
oob see represampling.factor.bootstrap
... additional arguments to be passed to partition.kmeans
data data.frame containing at least the columns specified by coords
coords vector of length 2 defining the variables in data that contain the x and y coordinates of sample locations
repetition numeric vector: cross-validation repetitions to be generated. Note that this is not the number of repetitions, but the indices of these repetitions. E.g., use repetition=c(1:100) to obtain the 'first' 100 repetitions, and repetition=c(101:200) to obtain a different set of 100 repetitions.
seed1 seed1+i is the random seed that will be used by set.seed in repetition i (i in repetition) to initialize the random number generator before sampling from the data set.

represampling.tile.bootstrap

Spatial block bootstrap using rectangular blocks

Description

represampling.tile.bootstrap performs a non-overlapping spatial block bootstrap by resampling at the level of rectangular partitions or 'tiles' generated by partition.tiles.

Usage

represampling.tile.bootstrap(data, coords = c("x", "y"),
   repetition = 1, nboot = -1, seed1 = NULL, oob = FALSE,
   ...)

Arguments

nboot see represampling.factor.bootstrap
oob see represampling.factor.bootstrap
... additional arguments to be passed to partition.tiles
data data.frame containing at least the columns specified by coords
resample.strat.uniform
draw stratified random sample

Description
resample.strat.uniform draws a stratified random sample (with or without replacement) from
the samples in data. Stratification is over the levels of data[,param$response]. The same number
of samples is drawn within each level.

Usage
resample.strat.uniform(data,
param = list(strat = "class", nstrat = Inf, replace = FALSE))

Arguments
data a data.frame, rows represent samples
param a list with the following components: strat is either the name of a factor vari-
able in data that defines the stratification levels, or a vector of type factor and
length nrow(data); n is a numeric value specifying the size of the subsample;
replace determines if sampling is with or without replacement

Details
If param$replace=FALSE, a subsample of size min(param$n,nrow(data)) will be drawn from
data. If param$replace=TRUE, the size of the subsample is param$n.

Value
a data.frame containing a subset of the rows of data.

See Also
resample.uniform, sample
resample.unifrom

Examples

```r
data(ecuador) # Muenchow et al. (2012), see ?ecuador
resample.strat.uniform(ecuador, param = list(strat = "slides", nstrat = 100))
nrow(d) # == 200
sum(d$slides == "TRUE") # == 100
```

---

**resample.unifrom**  
*Draw uniform random (sub)samples*

**Description**

`resample.unifrom` draws a random (sub)sample (with or without replacement) from the samples in `data`.

**Usage**

```r
resample.unifrom(data, param = list(n = Inf, replace = FALSE))
```

**Arguments**

- `data` a `data.frame`, rows represent samples
- `param` a list with the following components:  
  - `n` is a numeric value specifying the size of the subsample;  
  - `replace` determines if sampling is with or without replacement

**Details**

If `param$replace`=FALSE, a subsample of size `min(param$n, nrow(data))` will be drawn from `data`. If `param$replace`=TRUE, the size of the subsample is `param$n`.

**Value**

A `data.frame` containing a subset of the rows of `data`.

**See Also**

`resample.strat.unifrom`, `sample`

**Examples**

```r
data(ecuador) # Muenchow et al. (2012), see ?ecuador
resample.unifrom(ecuador, param = list(strat = "slides", n = 200))
nrow(d) # == 200
sum(d$slides == "TRUE")
```
Perform spatial error estimation and variable importance assessment

Description

sperrorest is a flexible interface for multiple types of spatial and non-spatial cross-validation and bootstrap error estimation and permutation-based assessment of spatial variable importance.

Usage

sperrorest(formula, data, coords = c("x", "y"),
  model.fun, model.args = list(), pred.fun = NULL,
  pred.args = list(), smp.fun = partition.loo,
  smp.args = list(), train.fun = NULL,
  train.param = NULL, test.fun = NULL, test.param = NULL,
  err.fun = err.default, err.unpooled = TRUE,
  err.pooled = FALSE, err.train = TRUE,
  imp.variables = NULL, imp.permutations = 1000,
  importance = !is.null(imp.variables), distance = FALSE,
  do.gc = 1, do.try = FALSE, silent = FALSE, ...)

Arguments

data a data.frame with predictor and response variables. Training and test samples will be drawn from this data set by train.fun and test.fun, respectively.

formula A formula specifying the variables used by the model. Only simple formulas without interactions or nonlinear terms should be used, e.g. \( y \sim x + x^2 + x^3 \) but not \( y \sim x + x^2 + \log(x)^3 \). Formulas involving interaction and nonlinear terms may possibly work for error estimation but not for variable importance assessment, but should be used with caution.

coords vector of length 2 defining the variables in data that contain the x and y coordinates of sample locations

model.fun Function that fits a predictive model, such as glm or rpart. The function must accept at least two arguments, the first one being a formula and the second a data.frame with the learning sample.

model.args Arguments to be passed to model.fun (in addition to the formula and data argument, which are provided by sperrorest)

pred.fun Prediction function for a fitted model object created by model. Must accept at least two arguments, the fitted object and a data.frame newdata with data on which to predict the outcome.

pred.args (optional) Arguments to pred.fun (in addition to the fitted model object and the newdata argument, which are provided by sperrorest)

smp.fun A function for sampling training and test sets from data. E.g., partition.kmeans for spatial cross-validation using spatial k-means clustering.

smp.args (optional) Arguments to be passed to est.fun
train.fun (optional) A function for resampling or subsampling the training sample in order to achieve, e.g., uniform sample sizes on all training sets, or maintaining a certain ratio of positives and negatives in training sets. E.g., `resample.uniform` or `resample.strat.uniform`

test.fun (optional) Like train.fun but for the test set.

test.param (optional) Arguments to be passed to test.fun

err.fun A function that calculates selected error measures from the known responses in data and the model predictions delivered by pred.fun. E.g., `err.default` (the default). See example and details below.

err.unpooled logical (default: TRUE): calculate error measures on each fold within a resampling repetition

err.pooled logical (default: FALSE): calculate error measures based on the pooled predictions of all folds within a resampling repetition

err.train logical (default: TRUE): calculate error measures on the training set (in addition to the test set estimation)

imp.variables (optional; used if importance=TRUE) Variables for which permutation-based variable importance assessment is performed. If importance=TRUE and imp.variables is NULL, all variables in formula will be used.

imp.permutations (optional; used if importance=TRUE) Number of permutations used for variable importance assessment.

importance logical: perform permutation-based variable importance assessment?

... currently not used

distance logical (default: FALSE): if TRUE, calculate mean nearest-neighbour distances from test samples to training samples using `add.distance.represampling`

do.gc numeric (default: 1): defines frequency of memory garbage collection by calling gc; if <1, no garbage collection; if >=1, run a gc() after each repetition; if >=2, after each fold

do.try logical (default: FALSE): if TRUE [untested!!], use try to robustify calls to model.fun and err.fun; use with caution!

silent If TRUE, show progress on console (in Windows Rgui, disable 'Buffered output' in 'Misc' menu)

Value

A list (object of class sperrorest) with (up to) four components:

- **error** a sperroresterror object containing predictive performances at the fold level
- **represampling** a represampling object
- **pooled.error** a sperrorestpoolederror object containing predictive performances at the repetition level
- **importance** a sperrorestimportance object containing permutation-based variable importances at the fold level
An object of class `sperrorest`, i.e. a list with components `error` (of class `sperroresterror`), `represampling` (of class `represampling`), `pooled.error` (of class `sperrorestpoolederror`) and `importance` (of class `sperrorestimportance`).

**Note**

To do: (1) Parallelize the code; (2) Optionally save fitted models, training and test samples in the results object; (3) Optionally save intermediate results in some file, and enable the function to continue an interrupted sperrorest call where it was interrupted. (3) Optionally have sperrorest dump the result of each repetition into a file, and to skip repetitions for which a file already exists. (4) Save sperrorest version number in results object.

**References**


**See Also**

`ipred`

**Examples**

```r
data(ecuador) # Muenchow et al. (2012), see ?ecuador
fo = slides ~ dem + slope + hcurv + vcurv +
     log.carea + cslope

# Example of a classification tree fitted to this data:
library(rpart)
ctrl = rpart.control(cp = 0.005) # show the effects of overfitting
fit = rpart(fo, data = ecuador, control = ctrl)
par(xpd = TRUE)
plot(fit, compress = TRUE, main = "Stoyan's landslide data set")
text(fit, use.n = TRUE)

# Non-spatial 5-repeated 10-fold cross-validation:
```

---

**sperrorest**

An object of class `sperrorest`, i.e. a list with components `error` (of class `sperroresterror`), `represampling` (of class `represampling`), `pooled.error` (of class `sperrorestpoolederror`) and `importance` (of class `sperrorestimportance`).

**Note**

To do: (1) Parallelize the code; (2) Optionally save fitted models, training and test samples in the results object; (3) Optionally save intermediate results in some file, and enable the function to continue an interrupted sperrorest call where it was interrupted. (3) Optionally have sperrorest dump the result of each repetition into a file, and to skip repetitions for which a file already exists. (4) Save sperrorest version number in results object.

**References**


**See Also**

`ipred`

**Examples**

```r
data(ecuador) # Muenchow et al. (2012), see ?ecuador
fo = slides ~ dem + slope + hcurv + vcurv +
     log.carea + cslope

# Example of a classification tree fitted to this data:
library(rpart)
ctrl = rpart.control(cp = 0.005) # show the effects of overfitting
fit = rpart(fo, data = ecuador, control = ctrl)
par(xpd = TRUE)
plot(fit, compress = TRUE, main = "Stoyan's landslide data set")
text(fit, use.n = TRUE)

# Non-spatial 5-repeated 10-fold cross-validation:
```
mypred.rpart = function(object, newdata) predict(object, newdata)[,2]
nspres = sperrorest(data = ecuador, formula = fo,
    model.fun = rpart, model.args = list(control = ctrl),
    pred.fun = mypred.rpart,
    smp.fun = partition.cv, smp.args = list(repetition=1:5, nfold=10))
summary(nspres$error)
summary(nspres$represampling)
plot(nspres$represampling, ecuador)

# Spatial 5-repeated 10-fold spatial cross-validation:
spres = sperrorest(data = ecuador, formula = fo,
    model.fun = rpart, model.args = list(control = ctrl),
    pred.fun = mypred.rpart,
    smp.fun = partition.kmeans, smp.args = list(repetition=1:5, nfold=10))
summary(spres$error)
summary(spres$represampling)
plot(spres$represampling, ecuador)

smry = data.frame(
    nonspat.training = unlist(summary(nspres$error, level=1)$train.auroc),
    nonspat.test = unlist(summary(nspres$error, level=1)$test.auroc),
    spatial.training = unlist(summary(spres$error, level=1)$train.auroc),
    spatial.test = unlist(summary(spres$error, level=1)$test.auroc))
boxplot(smry, col = c("red","red","red","green"),
    main = "Training vs. test, nonspatial vs. spatial",
    ylab = "Area under the ROC curve")

summary.nspres  Summary statistics for a resampling objects

Description

Calculates sample sizes of training and test sets within repetitions and folds of a resampling or represampling object.

Usage

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

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

Arguments

object  A resampling or represampling object.

...  currently ignored
summary.sperroresterror

Summarize error statistics obtained by sperrorest

Description

summary.sperroresterror calculates mean, standard deviation, median etc. of the calculated error measures at the specified level (overall, repetition, or fold). summary.sperrorestpoolederror does the same with the pooled error, at the overall or repetition level.

Usage

```r
## S3 method for class 'sperroresterror'
summary(object, level = 0,
         pooled = TRUE, na.rm = TRUE, ...)
```

Arguments

- **object** sperroresterror resp. sperrorestcombinederror error object calculated by sperrorest
- **level** Level at which errors are summarized: 0: overall; 1: repetition; 2: fold
- **pooled** If TRUE (default), mean and standard deviation etc are calculated between fold-level error estimates. If FALSE, apply first a weighted.mean among folds before calculating mean, standard deviation etc among repetitions. See also Details.
- **na.rm** Remove NA values? See mean etc.
- **...** additional arguments (currently ignored)

Details

Let’s use an example to explain the pooled argument. E.g., assume we are using 100-repeated 10-fold cross-validation. If pooled=TRUE (default), the mean and standard deviation calculated when summarizing at level=0 are calculated across the error estimates obtained for each of the 100*10 = 1000 folds. If pooled=FALSE, mean and standard deviation are calculated across the 100 repetitions, using the weighted average of the fold-level errors to calculate an error value for the entire sample. This will essentially not affect the mean value but of course the standard deviation of the error. pooled=FALSE is not recommended, it is mainly for testing purposes; when the test sets are small (as in leave-one-out cross-validation, in the extreme case), consider running sperrorest with err.pooled=TRUE and examine only the pooled.error component of its result.

Value

Depending on the level of aggregation, a list or data.frame with mean, and at level 0 also standard deviation, median and IQR of the error measures.
### summary.sperrorestimportance

**Summary and print methods for sperrorest results**

#### Description

Summary methods provide varying level of detail while print methods provide full details.

#### See Also

- sperrorest
tile.neighbors

Determine the names of neighbouring tiles in a rectangular pattern

Description

This based on "counting" up and down based on the tile name.

Usage

```
tile.neighbors(nm, tileset, iterate = 0, diagonal = FALSE)
```
Arguments

- **`nm`**: Character string or factor: name of a tile, e.g., "X4:Y6"
- **`tileset`**: Admissible tile names; if missing and `nm` is a factor variable, then `levels(nm)` is used as a default for `tileset`
- **`iterate`**: Internal - do not change default: to control behaviour in an interactive call to this function
- **`diagonal`**: If TRUE, diagonal neighbours are also considered neighbours

Value

Character string.
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