

Package ‘HPdclassifier’

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Type Package

Title Distributed classifiers for Big Data

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Depends R (>= 3.0.1), distributedR, randomForest

Description It provides distributed algorithms for learning classifiers. It is written based on the infrastructure created in HP-Labs for distributed computing in R.

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HPdclassifier-package
Distributed algorithms for classifiers

Description

HPdclassifier provides several distributed algorithms for classifiers. It is written based on the infrastructure created in HP-Labs for distributed computing in R.

Details

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1

Main Functions:

- hpdrandomForest: It is a distributed function for randomForest

Author(s)

Arash Fard <afard@vertica.com>

References

1. Using R for Iterative and Incremental Processing. Shivaram Venkataraman, Indrajit Roy, Alvin AuYoung, Rob Schreiber. HotCloud 2012, Boston, USA.

hpdrandomForest	<i>Distributed randomForest</i>
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Description

hpdrandomForest function runs randomForest function of randomForest package in a distributed fashion.

Description

hpdrandomForest calls several instances of randomForest distributed across a cluster system. Therefore, the master distributes the input data among all R-executors of the distributedR environment, and trees on different sub-sections of the forest are created simultaneously. At the end, all these trees are combined to result a single forest.

The interface of hpdrandomForest is similar to randomForest. Indeed it adds two arguments nExecutor and trace, and removes several other arguments do.trace, keep.inbag, proximity, and oob.prox. Its returned result is also completely compatible to the result of randomForest.

Usage

```
## S3 method for class 'formula'
hpdrandomForest(formula, data=NULL, ..., subset, na.action=na.fail,
                 nExecutor, trace=FALSE)

## Default S3 method:
hpdrandomForest(x, y=NULL, xtest=NULL, ytest=NULL, ntree=500,
  mtry=if (!is.null(y) && !is.factor(y) && !is.dframe(y))
    max(floor(ncol(x)/3), 1) else floor(sqrt(ncol(x))),
  replace=TRUE, classwt=NULL, cutoff, strata,
  sampsize = if (replace) nrow(x) else ceiling(.632*nrow(x)),
  nodesize = if (!is.null(y) && !is.factor(y) &&
    !is.dframe(y)) 5 else 1,
  maxnodes = NULL,
  importance=FALSE, localImp=FALSE, nPerm=1,
  norm.votes=TRUE,
  keep.forest=!is.null(y) && is.null(xtest),
  corr.bias=FALSE, nExecutor, trace=FALSE, ...)
```

Arguments

<code>data</code>	an optional data frame containing the variables in the model. By default the variables are taken from the environment which <code>hpdrandomForest</code> is called from.
<code>subset</code>	an index vector indicating which rows should be used. (NOTE: If given, this argument must be named.)
<code>na.action</code>	A function to specify the action to be taken if NAs are found. (NOTE: If given, this argument must be named.)
<code>formula</code>	a formula describing the model to be fitted.
<code>x</code>	when a data frame or a matrix of predictors assigned to <code>x</code> , its size should not be bigger than 64MB. For bigger datasets, <code>darray</code> or <code>dframe</code> should be used. <code>darray</code> is more memory efficient than <code>dframe</code> , but it does not support categorical data. Therefore, using <code>darray</code> is highly recommended when there is no categorical data.
<code>y</code>	a response vector. If a factor, classification is assumed, otherwise regression is assumed. If omitted, <code>hpdrandomForest</code> will run in unsupervised mode. When <code>x</code> is a distributed structure (<code>darray</code> or a <code>dframe</code>), <code>y</code> should be also a distributed structure with a single column.
<code>xtest</code>	a data frame or matrix (like <code>x</code>) containing predictors for the test set. When <code>x</code> is a <code>darray</code> or a <code>dframe</code> , it should be of the same type.
<code>ytest</code>	response for the test set. Its type should be consistent with <code>y</code> . Moreover, it should have a single column.
<code>ntree</code>	Number of trees to grow. This should not be set to too small a number, to ensure that every input row gets predicted at least a few times.
<code>mtry</code>	Number of variables randomly sampled as candidates at each split. Note that the default values are different for classification (\sqrt{p} where p is number of variables in <code>x</code>) and regression ($p/3$)
<code>replace</code>	Should sampling of cases be done with or without replacement?
<code>classwt</code>	Priors of the classes. Need not add up to one. Ignored for regression.
<code>cutoff</code>	(Classification only) A vector of length equal to number of classes. The 'winning' class for an observation is the one with the maximum ratio of proportion of votes to cutoff. Default is $1/k$ where k is the number of classes (i.e., majority vote wins).
<code>strata</code>	A (factor) variable that is used for stratified sampling.
<code>sampsize</code>	Size(s) of sample to draw. For classification, if <code>sampsize</code> is a vector of the length the number of strata, then sampling is stratified by strata, and the elements of <code>sampsize</code> indicate the numbers to be drawn from the strata.
<code>nodesize</code>	Minimum size of terminal nodes. Setting this number larger causes smaller trees to be grown (and thus take less time). Note that the default values are different for classification (1) and regression (5).
<code>maxnodes</code>	Maximum number of terminal nodes trees in the forest can have. If not given, trees are grown to the maximum possible (subject to limits by <code>nodesize</code>). If set larger than maximum possible, a warning is issued.
<code>importance</code>	Should importance of predictors be assessed?
<code>localImp</code>	Should casewise importance measure be computed? (Setting this to <code>TRUE</code> will override <code>importance</code> .)

nPerm	Number of times the OOB data are permuted per tree for assessing variable importance. Number larger than 1 gives slightly more stable estimate, but not very effective. Currently only implemented for regression.
norm.votes	If TRUE (default), the final result of votes are expressed as fractions. If FALSE, raw vote counts are returned (useful for combining results from different runs). Ignored for regression.
keep.forest	If set to FALSE, the forest will not be retained in the output object. If xtest is given, defaults to FALSE.
corr.bias	perform bias correction for regression? Note: Experimental. Use at your own risk.
...	optional parameters to be passed to the low level function.
nExecutor	a positive integer number indicating the number of tasks for running the function. To have optimal performance, it is recommended to have this number smaller than the number of R-executors in the distributedR environment. It cannot be bigger than ntree.
trace	when this argument is true, intermediate steps of the progress are displayed.

Value

An object of class `randomForest`. The result is consistent with the result of the `combine` function in `randomForest` package.

Note

When `ntree` is not big enough in comparison to `nExecutor`, some of the returned predicted values may become NULL.

Returned values for `err.rate`, `votes`, and `oob.times` are valid only for classification type.

Author(s)

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References

Breiman, L. (2001), *Random Forests*, Machine Learning 45(1), 5-32.

Breiman, L (2002), "Manual On Setting Up, Using, And Understanding Random Forests V3.1", http://oz.berkeley.edu/users/breiman/Using_random_forests_V3.1.pdf.

Random Forests V4.6-7, <http://cran.r-project.org/web/packages/randomForest/randomForest.pdf>.

Examples

```
## Not run:

library(HPdclassifier)
distributedR_start()
drs <- distributedR_status()
nparts <- sum(drs$Ins)

## Classification:
##data(iris)
iris.rf <- hpdrandomForest(Species ~ ., data=iris, importance=TRUE,
```

```

                                nExecutor=nparts)
print(iris.rf)

## The `unsupervised' case:
iris.urf <- hpdrandomForest(iris[, -5], nExecutor=nparts)
MDSplot(iris.urf, iris$Species)

## stratified sampling: draw 20, 30, and 20 of the species to grow each tree.
(iris.rf2 <- hpdrandomForest(iris[1:4], iris$Species,
                             sampsize=c(20, 30, 20), nExecutor=nparts))

## Regression:
## data(airquality)
ozone.rf <- hpdrandomForest(Ozone ~ ., data=airquality, mtry=3,
                             importance=TRUE, na.action=na.omit, nExecutor=nparts)
print(ozone.rf)
## Show "importance" of variables: higher value mean more important:
round(importance(ozone.rf), 2)

## "x" can be a matrix instead of a data frame:
x <- matrix(runif(5e2), 100)
y <- gl(2, 50)
(myrf <- hpdrandomForest(x, y, nExecutor=nparts))
(predict(myrf, x))

## "complicated" formula:
(swiss.rf <- hpdrandomForest(sqrt(Fertility) ~ . - Catholic + I(Catholic < 50),
                             data=swiss, nExecutor=nparts))
(predict(swiss.rf, swiss))
## Test use of 32-level factor as a predictor:
x <- data.frame(x1=gl(32, 5), x2=runif(160), y=rnorm(160))
(rfl <- hpdrandomForest(x[-3], x[[3]], ntree=10, nExecutor=nparts))

## Grow no more than 4 nodes per tree:
(treesize(hpdrandomForest(Species ~ ., data=iris, maxnodes=4, ntree=30,
                           nExecutor=nparts)))

distributedR_shutdown()

## End(Not run)

```

predictHPdRF

distributed predict method for applying a random forest objects on a darray or a dframe

Description

Prediction of distributed test data using random forest.

Usage

```
predictHPdRF(object, newdata, trace=FALSE)
```

Arguments

object	an object of class <code>randomForest</code> , as that created by the function <code>randomForest</code> or <code>hpdrandomForest</code> .
newdata	a darray or a dframe containing new data. darray is highly recommended when there is no categorial data

Value

It returns a darray or a dframe of predicted classes. The type of returned value will be darray unless there is any categorial data.

Author(s)

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References

Breiman, L. (2001), *Random Forests*, Machine Learning 45(1), 5-32.

See Also

[hpdrandomForest](#)

Examples

```
## Not run:
# example for darray
library(HPdclassifier)
distributedR_start()
drs <- distributedR_status()
nparts <- sum(drs$Ins)

nSamples <- 100
nAttributes <- 5
nSpllits <- 1

dax <- darray(c(nSamples,nAttributes), c(round(nSamples/nSpllits),nAttributes))
day <- darray(c(nSamples,1), c(round(nSamples/nSpllits),1))

foreach(i, 1:npartitions(dax), function(x=splits(dax,i),y=splits(day,i),id=i){
  x <- matrix(runif(nrow(x)*ncol(x)), nrow(x),ncol(x))
  y <- matrix(runif(nrow(y)), nrow(y), 1)
  update(x)
  update(y)
})

(myrf1 <- hpdrandomForest(dax, day, nExecutor=nparts))
dp <- predictHPdRF(myrf1, dax)

# example for dframe
nSamples <- 100
nAttributes <- 5
nSpllits <- 4

dfx <- dframe(c(nSamples,nAttributes), c(round(nSamples/nSpllits),nAttributes))
```

```
dfy <- dframe(c(nSamples,1), c(round(nSamples/nSpllits),1))
blockSize <- dfx@blocks[1]

foreach(i, 1:npartitions(dfx), function(xi=splits(dfx,i),yi=splits(dfy,i),
  blockSize=blockSize,nAttributes=nAttributes){
  xi <- data.frame( matrix(runif(blockSize*nAttributes,1,10),
    nrow=blockSize,ncol=nAttributes) )
  yi <- data.frame( gl(2, blockSize/2) )
  update(xi)
  update(yi)
})

(myrf2 <- hpdrandomForest(dfx, dfy, nExecutor=nparts))
fp2 <- predictHPdRF(myrf2, dfx)

## End(Not run)
```

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