

Package ‘caretEnsemble’

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Description Functions for creating ensembles of caret models: caretList() and caretStack(). caretList() is a convenience function for fitting multiple caret::train() models to the same dataset. caretStack() will make linear or non-linear combinations of these models, using a caret::train() model as a meta-model.

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add_cross_group_stats	<i>Add cross-group statistics to the importance table</i>
-----------------------	---

Description

Classifies each variable as either "Original" or "New" and optionally appends a summary statistic (mean, sum, or max) of the opposite group. The added statistic is flagged with 'is_stat = TRUE' and can be used for plotting a reference bar.

Usage

```
add_cross_group_stats(imp_dt, original_features, stat_type = NULL)
```

Arguments

imp_dt	data.table from 'prepare_importance'.
original_features	Character vector of original features.
stat_type	Character string: "mean", "sum", "max", or NULL.

Value

List with 'imp_original' and 'imp_new' data.tables.

as.caretList	<i>Convert object to caretList object</i>
--------------	---

Description

Converts object into a caretList

Usage

```
as.caretList(object)
```

Arguments

object	R Object
--------	----------

Value

a `caretList` object

as.caretList.default *Convert object to caretList object - For Future Use*

Description

Converts object into a caretList - For Future Use

Usage

```
## Default S3 method:  
as.caretList(object)
```

Arguments

object R object

Value

NA

as.caretList.list *Convert list to caretList*

Description

Converts list to caretList

Usage

```
## S3 method for class 'list'  
as.caretList(object)
```

Arguments

object list of caret models

Value

a `caretList` object

autoplot.caretStack	<i>Convenience function for more in-depth diagnostic plots of caretStack objects</i>
---------------------	--

Description

This function provides a more robust series of diagnostic plots for a caretEnsemble object.

Usage

```
## S3 method for class 'caretStack'  
autoplot(object, training_data = NULL, xvars = NULL, show_class_id = 2L, ...)
```

Arguments

object	a caretStack object
training_data	The data used to train the ensemble. Required if xvars is not NULL Must be in the same row order as when the models were trained.
xvars	a vector of the names of x variables to plot against residuals
show_class_id	For classification only: which class level to show on the plot
...	ignored

Value

A grid of diagnostic plots. Top left is the range of the performance metric across each component model along with its standard deviation. Top right is the residuals from the ensembled model plotted against fitted values. Middle left is a bar graph of the weights of the component models. Middle right is the disagreement in the residuals of the component models (unweighted) across the fitted values. Bottom left and bottom right are the plots of the residuals against two random or user specified variables. Note that the ensemble must have been trained with savePredictions = "final", which is required to get residuals from the stack for the plot.

Examples

```
set.seed(42)  
data(models.reg)  
ens <- caretStack(models.reg[1:2], method = "lm")  
autoplot(ens)
```

c.caretList	<i>S3 definition for concatenating caretList</i>
-------------	--

Description

take N objects of class caretList and concatenate them into a larger object of class caretList for future ensembling

Usage

```
## S3 method for class 'caretList'  
c(...)
```

Arguments

... the objects of class caretList or train to bind into a caretList

Value

a `caretList` object

Examples

```
data(iris)  
model_list1 <- caretList(Sepal.Width ~ .,  
  data = iris,  
  tuneList = list(  
    lm = caretModelSpec(method = "lm")  
  )  
)  
  
model_list2 <- caretList(Sepal.Width ~ .,  
  data = iris, tuneLength = 1L,  
  tuneList = list(  
    rf = caretModelSpec(method = "rf")  
  )  
)  
  
bigList <- c(model_list1, model_list2)
```

c.train	<i>S3 definition for concatenating train objects</i>
---------	--

Description

take N objects of class train and concatenate into an object of class caretList for future ensembling

Usage

```
## S3 method for class 'train'  
c(...)
```

Arguments

... the objects of class train to bind into a caretList

Value

a [caretList](#) object

Examples

```
data(iris)  
model_lm <- caret::train(Sepal.Length ~ .,  
  data = iris,  
  method = "lm"  
)  
  
model_rf <- caret::train(Sepal.Length ~ .,  
  data = iris,  
  method = "rf",  
  tuneLength = 1L  
)  
  
model_list <- c(model_lm, model_rf)
```

caretEnsemble	<i>Combine several predictive models via weights</i>
---------------	--

Description

Find a greedy, positive only linear combination of several [train](#) objects

Functions for creating ensembles of caret models: caretList and caretStack

Usage

```
caretEnsemble(all.models, excluded_class_id = 0L, tuneLength = 1L, ...)
```

Arguments

<code>all.models</code>	an object of class <code>caretList</code>
<code>excluded_class_id</code>	The integer level to exclude from binary classification or multiclass problems. By default no classes are excluded, as the greedy optimizer requires all classes because it cannot use negative coefficients.
<code>tuneLength</code>	The size of the grid to search for tuning the model. Defaults to 1, as the only parameter to optimize is the number of iterations, and the default of 100 works well.
<code>...</code>	additional arguments to pass <code>caret::train</code>

Details

`greedyMSE` works well when you want an ensemble that will never be worse than any single model in the dataset. In the worst case scenario, it will select the single best model, if none of them can be ensembled to improve the overall score. It will also never assign any model a negative coefficient, which can help avoid unintuitive cases at prediction time (e.g. if the correlations between predictors breaks down on new data, negative coefficients can lead to bad results).

Value

a `caretEnsemble` object

Note

Every model in the "library" must be a separate `train` object. For example, if you wish to combine a random forests with several different values of `mtry`, you must build a model for each value of `mtry`. If you use several values of `mtry` in one `train` model, (e.g. `tuneGrid = expand.grid(.mtry=2:5)`), `caret` will select the best value of `mtry` before we get a chance to include it in the ensemble. By default, RMSE is used to ensemble regression models, and AUC is used to ensemble Classification models. This function does not currently support multi-class problems

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See Also

Useful links:

- <https://zachmayer.github.io/caretEnsemble/>
- <https://github.com/zachmayer/caretEnsemble>
- Report bugs at <https://github.com/zachmayer/caretEnsemble/issues>

Examples

```
set.seed(42)
models <- caretList(iris[1:50, 1:2], iris[1:50, 3], methodList = c("rpart", "rf"))
ens <- caretEnsemble(models)
summary(ens)
```

caretList

Create a list of several train models from the caret package

Description

Build a list of train objects suitable for ensembling using the [caretStack](#) function.

Usage

```
caretList(
  ...,
  trControl = NULL,
  methodList = NULL,
  tuneList = NULL,
  metric = NULL,
  continue_on_fail = FALSE,
  trim = TRUE,
  aggregate_resamples = TRUE
)
```

Arguments

...	arguments to pass to train . Don't use the formula interface, its slower and buggier compared to the X, y interface. Use a data.table for X. Particularly if you have a large dataset and/or many models, using a data.table will avoid unnecessary copies of your data and can save a lot of time and RAM. These arguments will determine which train method gets dispatched.
trControl	a trainControl object. If NULL, will use defaultControl.
methodList	optional, a character vector of caret models to ensemble. One of methodList or tuneList must be specified.
tuneList	optional, a NAMED list of caretModelSpec objects. This much more flexible than methodList and allows the specification of model-specific parameters (e.g. passing trace=FALSE to nnet)

metric a string, the metric to optimize for. If NULL, we will choose a good one.

continue_on_fail logical, should a valid caretList be returned that excludes models that fail, default is FALSE

trim logical should the train models be trimmed to save memory and speed up stacking

aggregate_resamples logical, whether to aggregate stacked predictions. Default is TRUE.

Value

A list of `train` objects. If the model fails to build, it is dropped from the list.

Examples

```
caretList(
  Sepal.Length ~ Sepal.Width,
  head(iris, 50),
  methodList = c("glm", "lm"),
  tuneList = list(
    nnet = caretModelSpec(method = "nnet", trace = FALSE, tuneLength = 1L)
  )
)
```

caretModelSpec	<i>Generate a specification for fitting a caret model</i>
----------------	---

Description

A caret model specification consists of 2 parts: a model (as a string) and the arguments to the train call for fitting that model

Usage

```
caretModelSpec(method = "rf", ...)
```

Arguments

method the modeling method to pass to `caret::train`

... Other arguments that will eventually be passed to `caret::train`

Value

a list of lists

Examples

```
caretModelSpec("rf", tuneLength = 5L, preProcess = "ica")
```

caretStack	<i>Combine several predictive models via stacking</i>
------------	---

Description

Stack several `train` models using a `train` model.

Usage

```
caretStack(
  all.models,
  new_X = NULL,
  new_y = NULL,
  metric = NULL,
  trControl = NULL,
  excluded_class_id = 1L,
  original_features = NULL,
  aggregate_resamples = TRUE,
  ...
)
```

Arguments

<code>all.models</code>	a <code>caretList</code> , or an object coercible to a <code>caretList</code> (such as a list of train objects)
<code>new_X</code>	Data to predict on for the <code>caretList</code> , prior to training the stack (for transfer learning). if <code>NULL</code> , the stacked predictions will be extracted from the <code>caretList</code> models.
<code>new_y</code>	The outcome variable to predict on for the <code>caretList</code> , prior to training the stack (for transfer learning). If <code>NULL</code> , will use the observed levels from the first model in the caret stack. If 0, will include all levels.
<code>metric</code>	the metric to use for grid search on the stacking model.
<code>trControl</code>	a <code>trainControl</code> object to use for training the ensemble model. If <code>NULL</code> , will use <code>defaultControl</code> .
<code>excluded_class_id</code>	The integer level to exclude from binary classification or multiclass problems.
<code>original_features</code>	a character vector of the names of the original features to include in the stack or <code>NULL</code> to not include any features. These features will be added to the stacked predictions from the models to train the ensemble model.
<code>aggregate_resamples</code>	logical, whether to aggregate resamples by keys. Default is <code>TRUE</code> .
<code>...</code>	additional arguments to pass to the stacking model

Details

Uses either transfer learning or stacking to stack models. Assumes that all models were trained on the same number of rows of data, with the same target values. The features, cross-validation strategies, and model types (class vs reg) may vary however. If your stack of models were trained with different number of rows, please provide `new_X` and `new_y` so the models can predict on a common set of data for stacking.

If your models were trained on different columns, you should use stacking.

If you have both differing rows and columns in your model set, you are out of luck. You need at least a common set of rows during training (for stacking) or a common set of columns at inference time for transfer learning.

Value

S3 `caretStack` object

References

Caruana, R., Niculescu-Mizil, A., Crew, G., & Ksikes, A. (2004). Ensemble Selection from Libraries of Models. <https://www.cs.cornell.edu/~caruana/ctp/ct.papers/caruana.icml04.icdm06long.pdf>

Examples

```
models <- caretList(
  x = iris[1:50, 1:2],
  y = iris[1:50, 3],
  methodList = c("rpart", "glm")
)
caretStack(models, method = "glm")
```

defaultControl

Construct a default train control for use with caretList

Description

Unlike `caret::trainControl`, this function defaults to 5 fold CV. CV is good for stacking, as every observation is in the test set exactly once. We use 5 instead of 10 to save compute time, as `caretList` is for fitting many models. We also construct explicit fold indexes and return the stacked predictions, which are needed for stacking. For classification models we return class probabilities.

Usage

```
defaultControl(
  target,
  method = "cv",
  number = 5L,
  savePredictions = "final",
```

```

    index = caret::createFolds(target, k = number, list = TRUE, returnTrain = TRUE),
    is_class = is.factor(target) || is.character(target),
    is_binary = length(unique(target)) == 2L,
    ...
  )

```

Arguments

target	the target variable.
method	the method to use for trainControl.
number	the number of folds to use.
savePredictions	the type of predictions to save.
index	the fold indexes to use.
is_class	logical, is this a classification or regression problem.
is_binary	logical, is this binary classification.
...	other arguments to pass to trainControl

defaultMetric	<i>Construct a default metric</i>
---------------	-----------------------------------

Description

Caret defaults to RMSE for classification and RMSE for regression. For classification, I would rather use ROC.

Usage

```
defaultMetric(is_class, is_binary)
```

Arguments

is_class	logical, is this a classification or regression problem.
is_binary	logical, is this binary classification.

dotplot.caretStack	<i>Comparison dotplot for a caretStack object</i>
--------------------	---

Description

This is a function to make a dotplot from a caretStack. It uses dotplot from the caret package on all the models in the ensemble, excluding the final ensemble model. At the moment, this function only works if the ensembling model has the same number of resamples as the component models.

Usage

```
## S3 method for class 'caretStack'
dotplot(x, ...)
```

Arguments

x	An object of class caretStack
...	passed to dotplot

Examples

```
set.seed(42)
models <- caretList(
  x = iris[1:100, 1:2],
  y = iris[1:100, 3],
  methodList = c("rpart", "glm")
)
meta_model <- caretStack(models, method = "lm")
lattice::dotplot(meta_model)
```

extractMetric	<i>Generic function to extract accuracy metrics from various model objects</i>
---------------	--

Description

A generic function to extract cross-validated accuracy metrics from model objects.

Usage

```
extractMetric(x, ...)
```

Arguments

x	An object from which to extract metrics. The specific method will be dispatched based on the class of x.
...	Additional arguments passed to the specific methods.

Value

A [data.table](#)

See Also

[extractMetric.train](#), [extractMetric.caretList](#), [extractMetric.caretStack](#)

`extractMetric.caretList`

Extract accuracy metrics from a [caretList](#) object

Description

Extract the cross-validated accuracy metrics from each model in a `caretList`.

Usage

```
## S3 method for class 'caretList'  
extractMetric(x, ...)
```

Arguments

<code>x</code>	a <code>caretList</code> object
<code>...</code>	passed to <code>extractMetric.train</code>

Value

A `data.table` with metrics from each model.

`extractMetric.caretStack`

Extract accuracy metrics from a [caretStack](#) object

Description

Extract the cross-validated accuracy metrics from the ensemble model and individual models in a `caretStack`.

Usage

```
## S3 method for class 'caretStack'  
extractMetric(x, ...)
```

Arguments

`x` a caretStack object
`...` passed to `extractMetric.train` and `extractMetric.caretList`

Value

A data.table with metrics from the ensemble model and individual models.

<code>extractMetric.train</code>	<i>Extract accuracy metrics from a <code>train</code> model</i>
----------------------------------	---

Description

Extract the cross-validated accuracy metrics and their SDs from caret.

Usage

```
## S3 method for class 'train'
extractMetric(x, metric = NULL, ...)
```

Arguments

`x` a train object
`metric` a character string representing the metric to extract.
`...` ignored If NULL, uses the metric that was used to train the model.

Value

A numeric representing the metric desired metric.

<code>greedyMSE</code>	<i>Greedy optimization for MSE</i>
------------------------	------------------------------------

Description

Greedy optimization for minimizing the mean squared error. Works for classification and regression.

Usage

```
greedyMSE(X, Y, max_iter = 100L)
```

Arguments

X	A numeric matrix of features.
Y	A numeric matrix of target values.
max_iter	An integer scalar of the maximum number of iterations.

Value

A list with components:

model_weights	A numeric matrix of model_weights.
RMSE	A numeric scalar of the root mean squared error.
max_iter	An integer scalar of the maximum number of iterations.

greedyMSE_caret	<i>caret interface for greedyMSE</i>
-----------------	--------------------------------------

Description

caret interface for greedyMSE. greedyMSE works well when you want an ensemble that will never be worse than any single predictor in the dataset. It does not use an intercept and it does not allow for negative coefficients. This makes it highly constrained and in general does not work well on standard classification and regression problems. However, it does work well in the case of: * The predictors are highly correlated with each other * The predictors are highly correlated with the model * You expect or want positive only coefficients In the worse case, this method will select one input and use that, but in many other cases it will return a positive, weighted average of the inputs. Since it never uses negative weights, you never get into a scenario where one model is weighted negative and on new data you get were predictions because a correlation changed. Since this model will always be a positive weighted average of the inputs, it will rarely do worse than the individual models on new data.

Usage

```
greedyMSE_caret()
```

permutationImportance	<i>Permutation Importance</i>
-----------------------	-------------------------------

Description

Permute each variable in a dataset and use the change in predictions to calculate the importance of each variable. Based on the scikit learn implementation of permutation importance: https://scikit-learn.org/stable/modules/permutation_importance.html. However, we don't compare to the target by a metric. We JUST look at the change in the model's predictions, as measured by MAE. (for classification, this is like using a Brier score). We shuffle each variable and recompute the predictions before and after the shuffle. The difference in MAE. is the importance of that variable. We normalize by computing the MAE of the shuffled original predictions as an upper bound on the MAE and divide by this value. So a variable that, when shuffled, caused predictions as bad as shuffling the output predictions, we know that variable is 100. Similarly, as with regular permutation importance, a variable that, when shuffled, gives the same MAE as the original model has an importance of 0.

This method cannot yield negative importances. It is merely a measure of how much the models uses the variable, and does not tell you which variables help or hurt generalization. Use the model's cross-validated metrics to assess generalization.

Usage

```
permutationImportance(model, newdata, normalize = TRUE)
```

Arguments

model	A train object from the caret package.
newdata	A data.frame of new data to use to compute importances. Can be the training data.
normalize	A logical indicating whether to normalize the importances to sum to one.

Value

A named numeric vector of variable importances.

plot.caretList	<i>Plot a caretList object</i>
----------------	--------------------------------

Description

This function plots the performance of each model in a caretList object.

Usage

```
## S3 method for class 'caretList'
plot(x, metric = NULL, ...)
```

Arguments

x	a caretList object
metric	which metric to plot
...	ignored

Value

A ggplot2 object

plot.caretStack	<i>Plot a caretStack object</i>
-----------------	---------------------------------

Description

This function plots the performance of each model in a caretList object.

Usage

```
## S3 method for class 'caretStack'
plot(x, metric = NULL, ...)
```

Arguments

x	a caretStack object
metric	which metric to plot. If NULL, will use the default metric used to train the model.
...	ignored

Value

a ggplot2 object

plot_group	<i>Plot a group of variable importances</i>
------------	---

Description

Generates a ggplot bar chart for a given set of variable importances.

Usage

```
plot_group(imp_dt, title, fill_colors, y_max)
```

Arguments

imp_dt	data.table with columns 'method', 'weight', 'is_stat'.
title	Title for the plot.
fill_colors	Named vector specifying colors for each fill group.
y_max	Numeric maximum for the y-axis.

Value

ggplot object.

plot_variable_importance

Plot Variable Importance from a caretStack Model

Description

This function plots the variable importance from a stacked ensemble model ('caretStack'), separating original features from new (engineered) features. It optionally includes cross-group summary statistics (mean, sum, or max) from one feature group into the other for visual reference. It returns a ggplot object; if both original and new features are present, the plot will contain two facets (original and new features) within the same figure. This is useful for diagnosing which group of features contributes more to the stacked model.

Usage

```
plot_variable_importance(stack_model, newdata, stat_type = NULL)
```

Arguments

stack_model	A caretStack model trained using caretEnsemble. It should have an attribute original_features listing the original input variables. If this attribute is missing, all variables are treated as "new".
newdata	A data frame containing the data used for calculating variable importance. Typically this should be the validation or test set.
stat_type	Optional character string indicating which summary statistic of the opposite group to include as a gray bar for reference. Must be one of "mean", "sum", or "max". If NULL, no statistic is shown. If invalid, an error is thrown.

Details

- Variable importance is computed using `caret::varImp`.
- If the model lacks the `original_features` attribute, all variables are considered new.
- Requires the packages: `data.table`, `ggplot2`, and `caret`.

Value

A ggplot object. If the model includes both original and new features, the plot will contain two facets ("Original Features" and "New Features"). If stat_type is provided, a gray bar appears in each plot representing the selected summary statistic from the opposite group (e.g., mean of new features shown in the original features plot).

predict.caretList	<i>Create a matrix of predictions for each of the models in a caretList</i>
-------------------	---

Description

Make a matrix of predictions from a list of caret models

Usage

```
## S3 method for class 'caretList'
predict(
  object,
  newdata = NULL,
  verbose = FALSE,
  excluded_class_id = 1L,
  aggregate_resamples = TRUE,
  ...
)
```

Arguments

object	an object of class caretList
newdata	New data for predictions. It can be NULL, but this is ill-advised.
verbose	Logical. If FALSE no progress bar is printed if TRUE a progress bar is shown. Default FALSE.
excluded_class_id	Integer. The class id to drop when predicting for multiclass
aggregate_resamples	logical, whether to aggregate resamples by keys. Default is TRUE.
...	Other arguments to pass to predict.train

predict.caretStack *Make predictions from a caretStack*

Description

Make predictions from a caretStack. This function passes the data to each function in turn to make a matrix of predictions, and then multiplies that matrix by the vector of weights to get a single, combined vector of predictions.

Usage

```
## S3 method for class 'caretStack'
predict(
  object,
  newdata = NULL,
  se = FALSE,
  level = 0.95,
  excluded_class_id = 0L,
  return_class_only = FALSE,
  verbose = FALSE,
  aggregate_resamples = TRUE,
  ...
)
```

Arguments

object	a caretStack to make predictions from.
newdata	a new dataframe to make predictions on
se	logical, should prediction errors be produced? Default is false.
level	tolerance/confidence level should be returned
excluded_class_id	Which class to exclude from predictions. Note that if the caretStack was trained with an excluded_class_id, that class is ALWAYS excluded from the predictions from the caretList of input models. excluded_class_id for predict.caretStack is for the final ensemble model. So different classes could be excluded from the caretList models and the final ensemble model.
return_class_only	a logical indicating whether to return only the class predictions as a factor. If TRUE, the return will be a factor rather than a data.table. This is a convenience function, and should not be widely used. For example if you have a downstream process that consumes the output of the model, you should have that process consume probabilities for each class. This will make it easier to change prediction probability thresholds if needed in the future.
verbose	a logical indicating whether to print progress
aggregate_resamples	logical, whether to aggregate resamples by keys. Default is TRUE.

... arguments to pass to `predict.train` for the ensemble model. Do not specify type here. For classification, type will always be prob, and for regression, type will always be raw.

Details

Prediction weights are defined as variable importance in the stacked caret model. This is not available for all cases such as where the library model predictions are transformed before being passed to the stacking model.

Value

a data.table of predictions

Examples

```
models <- caretList(
  x = iris[1:100, 1:2],
  y = iris[1:100, 3],
  methodList = c("rpart", "glm")
)
meta_model <- caretStack(models, method = "lm")
RMSE(predict(meta_model, iris[101:150, 1:2]), iris[101:150, 3])
```

predict.greedyMSE	<i>Predict method for greedyMSE</i>
-------------------	-------------------------------------

Description

Predict method for greedyMSE objects.

Usage

```
## S3 method for class 'greedyMSE'
predict(object, newdata, return_labels = FALSE, ...)
```

Arguments

object	A greedyMSE object.
newdata	A numeric matrix of new data.
return_labels	A logical scalar of whether to return labels.
...	Additional arguments. Ignored.

Value

A numeric matrix of predictions.

prepare_importance	<i>Prepare variable importance data.table from a caretStack</i>
--------------------	---

Description

Extracts variable importance from a 'caretStack' model and returns a 'data.table' with columns 'method' and 'weight' sorted by importance.

Usage

```
prepare_importance(stack_model, newdata)
```

Arguments

stack_model	A trained 'caretStack' model.
newdata	A data frame used for calculating variable importance.

Value

'data.table' with columns 'method' (variable names) and 'weight' (importance).

print.caretStack	<i>Print a caretStack object</i>
------------------	----------------------------------

Description

This is a function to print a caretStack.

Usage

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

Arguments

x	An object of class caretStack
...	ignored

Examples

```
models <- caretList(
  x = iris[1:100, 1:2],
  y = iris[1:100, 3],
  methodList = c("rpart", "glm")
)
meta_model <- caretStack(models, method = "lm")
print(meta_model)
```

print.greedyMSE	<i>Print method for greedyMSE</i>
-----------------	-----------------------------------

Description

Print method for greedyMSE objects.

Usage

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

Arguments

x	A greedyMSE object.
...	Additional arguments. Ignored.

print.summary.caretList	<i>Print a summary.caretList object</i>
-------------------------	---

Description

This is a function to print a summary.caretList

Usage

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

Arguments

x	An object of class summary.caretList
...	ignored

```
print.summary.caretStack
```

Print a summary.caretStack object

Description

This is a function to print a summary.caretStack.

Usage

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

Arguments

x	An object of class summary.caretStack
...	ignored

```
summary.caretList
```

Summarize a caretList

Description

This function summarizes the performance of each model in a caretList object.

Usage

```
## S3 method for class 'caretList'  
summary(object, metric = NULL, ...)
```

Arguments

object	a caretList object
metric	The metric to show. If NULL will use the metric used to train each model
...	passed to extractMetric

Value

A data.table with metrics from each model.

summary.caretStack	<i>Summarize a caretStack object</i>
--------------------	--------------------------------------

Description

This is a function to summarize a caretStack.

Usage

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

Arguments

object	An object of class caretStack
...	ignored

Examples

```
models <- caretList(  
  x = iris[1:100, 1:2],  
  y = iris[1:100, 3],  
  methodList = c("rpart", "glm")  
)  
meta_model <- caretStack(models, method = "lm")  
summary(meta_model)
```

tuneCheck	<i>Check that the tuning parameters list supplied by the user is valid</i>
-----------	--

Description

This function makes sure the tuning parameters passed by the user are valid and have the proper naming, etc.

Usage

```
tuneCheck(x)
```

Arguments

x	a list of user-supplied tuning parameters and methods
---	---

varImp.caretStack	<i>Variable importance for caretStack</i>
-------------------	---

Description

This is a function to extract variable importance from a caretStack.

Usage

```
## S3 method for class 'caretStack'
varImp(object, newdata = NULL, normalize = TRUE, ...)
```

Arguments

object	An object of class caretStack
newdata	the data to use for computing importance. If NULL, will use the stacked predictions from the models
normalize	a logical indicating whether to normalize the importances to sum to one.
...	passed to predict.caretList

varImp.greedyMSE	<i>variable importance for a greedyMSE model</i>
------------------	--

Description

Variable importance for a greedyMSE model.

Usage

```
## S3 method for class 'greedyMSE'
varImp(object, ...)
```

Arguments

object	A greedyMSE object.
...	Additional arguments. Ignored.

wtd.sd	<i>Calculate a weighted standard deviation</i>
--------	--

Description

Used to weight deviations among ensembled model predictions

Usage

```
wtd.sd(x, w, na.rm = FALSE)
```

Arguments

x	a numeric vector
w	a vector of weights equal to length of x
na.rm	a logical indicating how to handle missing values, default = TRUE

[.caretList	<i>Index a caretList</i>
-------------	--------------------------

Description

Index a caret list to extract caret models into a new caretList object

Usage

```
## S3 method for class 'caretList'  
object[index]
```

Arguments

object	an object of class caretList
index	selected index

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