Package ‘h2o’

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Version 3.36.0.3
Type  Package
Title  R Interface for the 'H2O' Scalable Machine Learning Platform
Date  2022-02-16
Description  R interface for 'H2O', the scalable open source machine learning platform that offers parallelized implementations of many supervised and unsupervised machine learning algorithms such as Generalized Linear Models (GLM), Gradient Boosting Machines (including XGBoost), Random Forests, Deep Neural Networks (Deep Learning), Stacked Ensembles, Naive Bayes, Generalized Additive Models (GAM), ANOVA GLM, Cox Proportional Hazards, K-Means, PCA, ModelSelection, Word2Vec, as well as a fully automatic machine learning algorithm (H2O AutoML).
License  Apache License (== 2.0)
URL  https://github.com/h2oai/h2o-3
BugReports  https://h2oai.atlassian.net/projects/PUBDEV
NeedsCompilation  no
SystemRequirements  Java (>= 8, <= 17)
Depends  R (>= 2.13.0), methods, stats
Imports  graphics, tools, utils, RCurl, jsonlite
Suggests  ggplot2 (>= 3.3.0), mlbench, Matrix, slam, bit64 (>= 0.9.7), data.table (>= 1.9.8), rgl (>= 0.100.19), plot3Drgl (>= 1.0.1), survival, DT, IRdisplay, htmltools, plotly, repr
R topics documented:

- `locate.R`
- `grid.R`
- `segment.R`
- `predict.R`
- `tf-idf.R`
- `rulefit.R`
- `explain.R`
- `permutation_varimp.R`
- `infogram.R`
- `zzz.R`
- `upliftrandomforest.R`

**RoxygenNote 7.1.2**

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**Repository** CRAN

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This is a package for running H2O via its REST API from within R. To communicate with a H2O instance, the version of the R package must match the version of H2O. When connecting to a new H2O cluster, it is necessary to re-run the initializer.

Details

Package: h2o
Type: Package
Version: 3.36.0.3
Branch: rel-zorn
Date: Wed Feb 16 17:54:16 UTC 2022
License: Apache License (== 2.0)
Depends: R (>= 2.13.0), RCurl, jsonlite, statmod, tools, methods, utils
H2O is the scalable open source machine learning platform that offers parallelized implementations of many supervised and unsupervised machine learning algorithms such as Generalized Linear Models (GLM), Gradient Boosting Machines (including XGBoost), Random Forests, Deep Neural Networks (Deep Learning), Stacked Ensembles, Naïve Bayes, Generalized Additive Models (GAM), ANOVA GLM, Maximum R GLM (maxrglm), Cox Proportional Hazards, K-Means, PCA, Word2Vec, as well as a fully automatic machine learning algorithm (H2O AutoML). As an example, to run GLM, call \texttt{h2o.glm} with the H2O parsed data and parameters (response variable, error distribution, etc.) as arguments.

This package enables the use of the H2O machine learning platform commands in R. To use H2O from R, you must start or connect to the "H2O cluster", the term we use to describe the backend H2O Java engine. To run H2O on your local machine, call \texttt{h2o.init} without any arguments, and H2O will be automatically launched at localhost:54321, where the IP is "127.0.0.1" and the port is 54321. If you have the H2O cluster running on a remote machine (e.g. AWS EC2), you must provide the IP and port of the remote machine as arguments to the \texttt{h2o.init} call.

Note that no actual data is stored in the R workspace; and no actual work is carried out by R. R only saves the named objects, which uniquely identify the data set, model, etc on the server. When the user makes a request, R queries the server via the REST API, which returns a JSON file with the relevant information that R then displays in the console.

\section*{Author(s)}
Maintainer: Erin LeDell <erin@h2o.ai>

\section*{References}
- H2O.ai Homepage
- H2O User Guide
- H2O on GitHub

\section*{Description}
Append a \texttt{<key,value>} pair to a list.

\section*{Usage}
\texttt{.addParm(parms, k, v)}

\section*{Arguments}
\begin{itemize}
  \item \textbf{parms} \hspace{1cm} a list to add the \texttt{<k,v>} pair to
  \item \textbf{k} \hspace{1cm} a key, typically the name of some algorithm parameter
  \item \textbf{v} \hspace{1cm} a value, the value of the algorithm parameter
\end{itemize}
Details

Contained here are a set of helper methods that perform type checking on the value passed in.

---

**.check_for_ggplot2**  
*Stop with a user friendly message if a user is missing the ggplot2 package or has an old version of it.*

---

Description

Stop with a user friendly message if a user is missing the ggplot2 package or has an old version of it.

Usage

`.check_for_ggplot2(version = "3.0.0")`

Arguments

- `version`: minimal required ggplot2 version

---

**.collapse**  
*Helper Collapse Function*

---

Description

Collapse a character vector into a `,`-sep array of the form: [thing1,thing2,...]

Usage

`.collapse(v)`

Arguments

- `v`: Character vector.
.consolidate_varimps  Consolidate variable importances

Description
Consolidation works in the following way: 1. if varimp variable is in x => add it to consolidated_varimps 2. for all remaining varimp variables: 1. find the longest prefix of varimp variable that is in x and add it to the consolidated varimp 2. if there was no match, throw an error 3. normalize the consolidated_varimps so they sum up to 1

Usage
.consolidate_varimps(model)

Arguments

model  H2OModel

Value
sorted named vector

.create_leaderboard  Create a leaderboard like data frame for models

Description
Create a leaderboard like data frame for models

Usage
.create_leaderboard(models_info, leaderboard_frame, top_n = 20)

Arguments

models_info  H2OAutoML object or list of models
leaderboard_frame  when provided with list of models, use this frame to calculate metrics
top_n  create leaderboard with just top_n models

Value
a data.frame
`.find_appropriate_column_name`  
*Tries to match a fuzzy_col_name with a column name that exists in cols.*

**Description**  
Tries to match a fuzzy_col_name with a column name that exists in cols.

**Usage**  
`.find_appropriate_column_name(fuzzy_col_name, cols)`

**Arguments**  
- `fuzzy_col_name` a name to be decoded  
- `cols` vector of columns that contain all possible column names, i.e., decode fuzzy_col_name must be in cols

**Value**  
a correct column name

---

`.customized_call`  
*A helper function that makes it easier to override/add params in a function call.*

**Description**  
A helper function that makes it easier to override/add params in a function call.

**Usage**  
`.customized_call(fun, ..., overridable_defaults = NULL, overrides = NULL)`

**Arguments**  
- `fun` Function to be called  
- `...` Parameters that can’t be overridden  
- `overridable_defaults` List of parameters and values that can be overridden  
- `overrides` Parameters to add/override.

**Value**  
result of fun
**.get_algorithm**  
*Get the algorithm used by the model_or_model_id*

**Description**  
Get the algorithm used by the model_or_model_id

**Usage**  
\[.get_algorithm(model_or_model_id, treat_xrt_as_algorithm = FALSE)\]

**Arguments**
- **model_or_model_id**  
  Model object or a string containing model id
- **treat_xrt_as_algorithm**  
  Try to find out if a model is XRT and if so report it as xrt

**Value**  
algorithm name

**.get_domain_mapping**  
*Get a mapping between columns and their domains*

**Description**  
Get a mapping between columns and their domains

**Usage**  
\[.get_domain_mapping(model)\]

**Arguments**
- **model**  
  an h2o model

**Value**  
list containing a mapping from column to its domains (levels)
`.get_feature_count`  
*Get feature count sorted by the count descending.*

**Description**
Get feature count sorted by the count descending.

**Usage**
`.get_feature_count(column)`

**Arguments**
- `column`  
  H2OFrame column

**Value**
named vector with feature counts

---

`.get_first_of_family`  
*Get first of family models*

**Description**
Get first of family models

**Usage**
`.get_first_of_family(models, all_stackedensembles = FALSE)`

**Arguments**
- `models`  
  models or model ids
- `all_stackedensembles`  
  if TRUE, select all stacked ensembles
.h2o.doGET

Just like doRawGET but fills in the default h2oRestApiVersion if none is provided

Description
Just like doRawGET but fills in the default h2oRestApiVersion if none is provided

Usage
.h2o.doGET(h2oRestApiVersion, urlSuffix, parms, ...)

Arguments

- **h2oRestApiVersion**
  
  (Optional) A version number to prefix to the urlSuffix. If no version is provided, a default version is chosen for you.

- **urlSuffix**
  
  The partial URL suffix to add to the calculated base URL for the instance

- **parms**
  
  (Optional) Parameters to include in the request

- **...**
  
  (Optional) Additional parameters.

Value
A list object as described above

.h2o.doPOST

Just like doRawPOST but fills in the default h2oRestApiVersion if none is provided

Description
Just like doRawPOST but fills in the default h2oRestApiVersion if none is provided

Usage
.h2o.doPOST(h2oRestApiVersion, urlSuffix, parms, ...)

Arguments

- **h2oRestApiVersion**
  
  (Optional) A version number to prefix to the urlSuffix. If no version is provided, a default version is chosen for you.

- **urlSuffix**
  
  The partial URL suffix to add to the calculated base URL for the instance

- **parms**
  
  (Optional) Parameters to include in the request

- **...**
  
  (Optional) Additional parameters.
.h2o.doRawGET

**Value**

A list object as described above

---

**.h2o.doRawGET**

*Perform a low-level HTTP GET operation on an H2O instance*

---

**Description**

Does not do any I/O level error checking. Caller must do its own validations. Does not modify the response payload in any way. Log the request and response if h2o.startLogging() has been called.

**Usage**

```r
.h2o.doRawGET(
  conn = h2o.getConnection(),
  h2oRestApiVersion,
  urlSuffix,
  parms,
  ...
)
```

**Arguments**

- `conn` *H2OConnection*
- `h2oRestApiVersion` *(Optional)* A version number to prefix to the `urlSuffix`. If no version is provided, the version prefix is skipped.
- `urlSuffix` The partial URL suffix to add to the calculated base URL for the instance
- `parms` *(Optional)* Parameters to include in the request
- `...` *(Optional)* Additional parameters.

**Details**

The return value is a list as follows: `$url` – Final calculated URL. `$postBody` – The body of the POST request from client to server. `$curlError` – TRUE if a socket-level error occurred. FALSE otherwise. `$curlErrorMessage` – If curlError is TRUE a message about the error. `$statusCode` – The HTTP status code. Usually 200 if the request succeeded. `$statusMessage` – A string describing the `statusCode`. `$payload` – The raw response payload as a character vector.

**Value**

A list object as described above
.h2o.doRawPOST

Perform a low-level HTTP POST operation on an H2O instance

Description

Does not do any I/O level error checking. Caller must do its own validations. Does not modify the response payload in any way. Log the request and response if h2o.startLogging() has been called.

Usage

.h2o.doRawPOST(
    conn = h2o.getConnection(),
    h2oRestApiVersion,
    urlSuffix,
    parms,
    fileUploadInfo,
    ...
)

Arguments

conn H2OConnection
h2oRestApiVersion
    (Optional) A version number to prefix to the urlSuffix. If no version is provided, the version prefix is skipped.
urlSuffix The partial URL suffix to add to the calculated base URL for the instance
parms
    (Optional) Parameters to include in the request
fileUploadInfo
    (Optional) Information to POST (NOTE: changes Content-type from XXX-www-url-encoded to multi-part). Use fileUpload(normalizePath("/path/to/file")).
...
    (Optional) Additional parameters.

Details

The return value is a list as follows: $url – Final calculated URL. $postBody – The body of the POST request from client to server. $curlError – TRUE if a socket-level error occurred. FALSE otherwise. $curlErrorMessage – If $curlError is TRUE a message about the error. $httpStatusCode – The HTTP status code. Usually 200 if the request succeeded. $httpStatusMessage – A string describing the httpStatusCode. $payload – The raw response payload as a character vector.

Value

A list object as described above
**h2o.doSafeGET**

Perform a safe (i.e. error-checked) HTTP GET request to an H2O cluster.

**Description**

This function validates that no CURL error occurred and that the HTTP response code is successful. If a failure occurred, then stop() is called with an error message. Since all necessary error checking is done inside this call, the valid payload is directly returned if the function successfully finishes without calling stop().

**Usage**

```r
.h2o.doSafeGET(h2oRestApiVersion, urlSuffix, parms, ...)
```

**Arguments**

- `h2oRestApiVersion` *(Optional)* A version number to prefix to the `urlSuffix`. If no version is provided, a default version is chosen for you.
- `urlSuffix` The partial URL suffix to add to the calculated base URL for the instance
- `parms` *(Optional)* Parameters to include in the request
- `...` *(Optional)* Additional parameters.

**Value**

The raw response payload as a character vector

---

**h2o.doSafePOST**

Perform a safe (i.e. error-checked) HTTP POST request to an H2O cluster.

**Description**

This function validates that no CURL error occurred and that the HTTP response code is successful. If a failure occurred, then stop() is called with an error message. Since all necessary error checking is done inside this call, the valid payload is directly returned if the function successfully finishes without calling stop().

**Usage**

```r
.h2o.doSafePOST(h2oRestApiVersion, urlSuffix, parms, fileUploadInfo, ...)
```
Arguments

- `h2oRestApiVersion` (Optional) A version number to prefix to the `urlSuffix`. If no version is provided, a default version is chosen for you.
- `urlSuffix` The partial URL suffix to add to the calculated base URL for the instance
- `parms` (Optional) Parameters to include in the request
- `fileUploadInfo` (Optional) Information to POST (NOTE: changes Content-type from `XXX-www-url-encoded` to multi-part). Use `fileUpload(normalizePath("/path/to/file"))`.
- `...` (Optional) Additional parameters.

Value

The raw response payload as a character vector

---

`.h2o.is_progress` **Check if Progress Bar is Enabled**

Description

Check if Progress Bar is Enabled

Usage

`.h2o.is_progress()`

---

`.h2o.locate` *Locate a file given the pattern `<bucket>/<path/to/file>` e.g. `h2o:::.h2o.locate("smalldata/iris/iris22.csv")` returns the absolute path to `iris22.csv`*

Description

Locate a file given the pattern `<bucket>/<path/to/file>` e.g. `h2o:::.h2o.locate("smalldata/iris/iris22.csv")` returns the absolute path to `iris22.csv`

Usage

`.h2o.locate(pathStub, root.parent = NULL)`

Arguments

- `pathStub` relative path
- `root.parent` search root directory
.h2o.perfect_auc

*Internal function that calculates a precise AUC from given probabilities and actual responses.*

Description

Note: The underlying implementation is not distributed and can only handle limited size of data. For internal use only.

Usage

```
.h2o.perfect_auc(probs, acts)
```

Arguments

- `probs` An H2OFrame holding vector of probabilities.
- `acts` An H2OFrame holding vector of actuals.

.h2o.primitives

*Map of operations known to H2O*

Description

Map of operations known to H2O

Usage

```
.h2o.primitives
```

Format

An object of class character of length 39.
Capabilities endpoints

Description
Capabilities endpoints

Usage
.h2o.__ALL_CAPABILITIES

Format
An object of class character of length 1.

Check H2O Server Health

Description
Warn if there are sick nodes.

Usage
.h2o.__checkConnectionHealth()

H2OFrame Manipulation

Description
H2OFrame Manipulation

Usage
.h2o.__CREATE_FRAME

Format
An object of class character of length 1.
Decryption Endpoints

Description
Decryption Endpoints

Usage
.h2o.__DECRYPTION_SETUP

Format
An object of class character of length 1.

Removal Endpoints

Description
Removal Endpoints

Usage
.h2o.__DKV

Format
An object of class character of length 1.

Export Files Endpoint Generator

Description
Export Files Endpoint Generator

Usage
.h2o.__EXPORT_FILES(frame)

Arguments
frame H2OFrame
Inspect/Summary Endpoints

**Description**
Inspect/Summary Endpoints

**Usage**
.h2o.__FRAMES

**Format**
An object of class character of length 1.

Import/Export Endpoints

**Description**
Import/Export Endpoints

**Usage**
.h2o.__IMPORT

**Format**
An object of class character of length 1.

Administrative Endpoints

**Description**
Administrative Endpoints

**Usage**
.h2o.__JOBS

**Format**
An object of class character of length 1.
.h2o.__LOGANDECHO  Log and Echo Endpoint

Description
Log and Echo Endpoint

Usage
.h2o.__LOGANDECHO

Format
An object of class character of length 1.

.h2o.__MODELS  Model Endpoint

Description
Model Endpoint

Usage
.h2o.__MODELS

Format
An object of class character of length 1.

.h2o.__MODEL_BUILDERS  Model Builder Endpoint Generator

Description
Model Builder Endpoint Generator

Usage
.h2o.__MODEL_BUILDERS(algo)

Arguments
algo  Canonical identifier of H2O algorithm.
**Model Metrics Endpoint**

**Description**

Model Metrics Endpoint

**Usage**

```
.h2o.__MODEL_METRICS(model, data)
```

**Arguments**

- `model`: H2OModel.
- `data`: H2OFrame.

**Parse Endpoints**

**Description**

Parse Endpoints

**Usage**

```
.h2o.__PARSE_SETUP
```

**Format**

An object of class character of length 1.

**Rapids Endpoint**

**Description**

Rapids Endpoint

**Usage**

```
.h2o.__RAPIDS
```

**Format**

An object of class character of length 1.
H2O Package Constants

Description

The API endpoints for interacting with H2O via REST are named here.

Usage

.h2o.__REST_API_VERSION

Format

An object of class integer of length 1.

Details

Additionally, environment variables for the H2O package are named here. Endpoint Version

Segment Models Builder Endpoint Generator

Description

Segment Models Builder Endpoint Generator

Usage

.h2o.__SEGMENT_MODELS_BUILDERS(algo)

Arguments

algo  Cannonical identifier of H2O algorithm.
Description

Word2Vec Endpoints

Usage

.h2o.__W2V_SYNONYMS

Format

An object of class character of length 1.

.has_varimp

Has the model variable importance?

Description

Has the model variable importance?

Usage

.has_varimp(model)

Arguments

model model or a string containing model id

Value

boolean
.interpretable

Description
Is the model considered to be interpretable, i.e., simple enough.

Usage
.interpretable(model)

Arguments
model model or a string containing model id

Value
boolean

.is_h2o_model

Description
Is the model an H2O model?

Usage
.is_h2o_model(model)

Arguments
model Either H2O model/model id => TRUE, or something else => FALSE

Value
boolean
### .is_h2o_tree_model

**Is the model a Tree-based H2O Model?**

**Description**

Is the model a Tree-based H2O Model?

**Usage**

```
.is_h2o_tree_model(model)
```

**Arguments**

- `model` Either tree-based H2O model/model id => TRUE, or something else => FALSE

**Value**

boolean

---

### .is_plotting_to_rnotebook

**Check if we are plotting in to r notebook.**

**Description**

Check if we are plotting in to r notebook.

**Usage**

```
.is_plotting_to_rnotebook()
```

**Value**

boolean
Enhance leaderboard with per-model predictions.

**Usage**

```
.leaderboard_for_row(models_info, newdata, row_index, top_n = 20)
```

**Arguments**

- `models_info`: models_info object
- `newdata`: H2OFrame
- `row_index`: index of the inspected row
- `top_n`: leaderboard will contain top_n models

**Value**

H2OFrame

Min-max normalization.

**Description**

Min-max normalization.

**Usage**

```
.min_max(col)
```

**Arguments**

- `col`: numeric vector

**Value**

normalized numeric vector
.model_ids

Get Model Ids

Description

When provided with list of models it will extract model ids. When provided with model ids it won’t change anything. Works for mixed list as well.

Usage

.model_ids(models)

Arguments

models list or vector of models/model_ids

Value

a vector of model_ids

.pkg.env

The H2O Package Environment

Description

The H2O Package Environment

Usage

.pkg.env

Format

An object of class environment of length 4.
.plot_varimp

Plot variable importances with ggplot2

Description

Plot variable importances with ggplot2

Usage

.plot_varimp(model, top_n = 10)

Arguments

model H2OModel
top_n Plot just top_n features

Value

list of variable importance, grouped variable importance, and variable importance plot

.process_models_or_automl

Do basic validation and transform object to a "standardized" list containing models, and their properties such as x, y, whether it is a (multinomial) classification or not etc.

Description

Do basic validation and transform object to a "standardized" list containing models, and their properties such as x, y, whether it is a (multinomial) classification or not etc.

Usage

.process_models_or_automl(
  object,
  newdata,
  require_single_model = FALSE,
  require_multiple_models = FALSE,
  top_n_from_AutoML = NA,
  only_with_varimp = FALSE,
  best_of_family = FALSE,
  require_newdata = TRUE
)
.shorten_model_ids

Arguments

object Can be a single model/model_id, vector of model_id, list of models, H2OAutoML object
newdata An H2OFrame with the same format as training frame
require_single_model If true, make sure we were provided only one model
require_multiple_models If true, make sure we were provided at least two models
top_n_from_AutoML If set, don’t return more than top_n models (applies only for AutoML object)
only_with_varimp If TRUE, return only models that have variable importance
best_of_family If TRUE, return only the best of family models; if FALSE return all models in object
require_newdata If TRUE, require newdata to be specified; otherwise allow NULL instead, this can be used when there is no need to know if the problem is (multinomial) classification.

Value

a list with the following names leader, is_automl, models, is_classification, is_multinomial_classification, x, y, model

Description

Shortens model ids if possible (iff there will be same amount of unique model_ids as before)

Usage

.shorten_model_ids(model_ids)

Arguments

model_ids character vector

Value

character vector
H2O <-> R Communication and Utility Methods

Description

Collected here are the various methods used by the h2o-R package to communicate with the H2O backend. There are methods for checking cluster health, polling, and inspecting objects in the H2O store.

Usage

`.skip_if_not_developer()`

Description

Convert to quantiles when provided with numeric vector. When `col` is a factor vector assign uniformly value between 0 and 1 to each level.

Usage

`.uniformize(col)`

Arguments

`col` vector

Value

vector with values between 0 and 1
.varimp

Get variable importance in a standardized way.

Description
Get variable importance in a standardized way.

Usage
.varimp(model)

Arguments
model H2OModel

Value
A named vector

.verify_dataxy

Used to verify data, x, y and turn into the appropriate things

Description
Used to verify data, x, y and turn into the appropriate things

Usage
.verify_dataxy(data, x, y, autoencoder = FALSE)

Arguments
data H2OFrame
x features
y response
autoencoder autoencoder flag
Description

Starting H2O For examples

Examples

```r
## Not run:
if (Sys.info()$sysname == "Darwin" && Sys.info()$release == '13.4.0') {
  quit(save = "no")
} else {
  h2o.init(nthreads = 2)
}
## End(Not run)
```

apply

Apply on H2O Datasets

Description

Method for apply on H2OFrame objects.

Usage

```r
apply(X, MARGIN, FUN, ...)
```

Arguments

- **X**: an H2OFrame object on which `apply` will operate.
- **MARGIN**: the vector on which the function will be applied over, either 1 for rows or 2 for columns.
- **FUN**: the function to be applied.
- **...**: optional arguments to `FUN`.

Value

Produces a new H2OFrame of the output of the applied function. The output is stored in H2O so that it can be used in subsequent H2O processes.

See Also

- `apply` for the base generic
Examples

## Not run:
library(h2o)
h2o.init()
iris_hf <- as.h2o(iris)
summary(apply(iris_hf, 2, sum))

## End(Not run)

as.character.H2OFrame  Convert an H2OFrame to a String

Description

Convert an H2OFrame to a String

Usage

## S3 method for class 'H2OFrame'
as.character(x, ...)

Arguments

x  An H2OFrame object
...
Further arguments to be passed from or to other methods.

Examples

## Not run:
library(h2o)
h2o.init()
pretrained <- as.h2o(data.frame(  
  C1 = c("a", "b"), C2 = c(0, 1), C3 = c(1, 0), C4 = c(0.2, 0.8),  
  stringsAsFactors = FALSE))
pretrained_w2v <- h2o.word2vec(pre_trained = pretrained, vec_size = 3)
words <- as.character(as.h2o(c("b", "a", "c", NA, "a")))
vecs <- h2o.transform(pretrained_w2v, words = words)

## End(Not run)
Converts parsed H2O data into an R data frame

Description

Downloads the H2O data and then scans it in to an R data frame.

Usage

## S3 method for class 'H2OFrame'
as.data.frame(x, ...)

Arguments

x An H2OFrame object.
... Further arguments to be passed down from other methods.

Details

Method as.data.frame.H2OFrame will use fread if data.table package is installed in required version.

See Also

use.package

Examples

## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
as.data.frame(prostate)

## End(Not run)
**Description**

Converts a collection of Segment Models to a data.frame

**Usage**

```r
## S3 method for class 'H2OSegmentModels'
as.data.frame(x, ...)
```

**Arguments**

- `x` Object of class `H2OSegmentModels`
- `...` Further arguments to be passed down from other methods.

**Value**

Returns data.frame with result of segment model training.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
iris_hf <- as.h2o(iris)
models <- h2o.train_segments(algorithm = "gbm",
                            segment_columns = "Species",
                            x = c(1:3), y = 4,
                            training_frame = iris_hf,
                            ntrees = 5,
                            max_depth = 4)
as.data.frame(models)

## End(Not run)
```

---

**as.factor**

*Convert H2O Data to Factors*

**Description**

Convert column/columns in the current frame to categoricals.
Usage

as.factor(x)

Arguments

x          a column from an H2OFrame data set.

See Also

as.factor.

Examples

## Not run:
library(h2o)
h2o.init()

    # Single column
cars <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/junit/cars_20mpg.csv"
df <- h2o.importFile(cars)
df["cylinders"] <- as.factor(df["cylinders"])
h2o.describe(df["cylinders"])

    # Multiple columns
cars <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/junit/cars_20mpg.csv"
df <- h2o.importFile(cars)
df["cylinders","economy_20mpg"] <- as.factor(df["cylinders","economy_20mpg"])
h2o.describe(df["cylinders","economy_20mpg"])

## End(Not run)

as.h2o

Create H2OFrame

Description

Import R object to the H2O cluster.

Usage

as.h2o(x, destination_frame = "", ...)  

## Default S3 method:
as.h2o(x, destination_frame = "", ...)

## S3 method for class 'H2OFrame'
as.h2o(x, destination_frame = "", ...)

## S3 method for class 'data.frame'
as.h2o (x, destination_frame = "", use_datatable = TRUE, ...)

## S3 method for class 'Matrix'
as.h2o(x, destination_frame = "", use_datatable = TRUE, ...)

Arguments

x       An R object.
destination_frame A string with the desired name for the H2OFrame
...      arguments passed to method arguments.
use_datatable allow usage of data.table

Details

Method as.h2o.data.frame will use fwrite if data.table package is installed in required version.
To speedup execution time for large sparse matrices, use h2o datatable. Make sure you have in-
stalled and imported data.table and slam packages. Turn on h2o datatable by options("h2o.use.data.table"=TRUE)

References

https://www.h2o.ai/blog/fast-csv-writing-for-r/

See Also

use.package

Examples

## Not run:
library(h2o)
h2o.init()
iris_hf <- as.h2o(iris)
euro_hf <- as.h2o(euro)
letters_hf <- as.h2o(letters)
state_hf <- as.h2o(state.x77)
iris_hf_2 <- as.h2o(iris_hf)
stopifnot(is.h2o(iris_hf), dim(iris_hf) == dim(iris),
is.h2o(euro_hf), dim(euro_hf) == c(length(euro), 1L),
is.h2o(letters_hf), dim(letters_hf) == c(length(letters), 1L),
is.h2o(state_hf), dim(state_hf) == dim(state.x77),
is.h2o(iris_hf_2), dim(iris_hf_2) == dim(iris_hf))
if (requireNamespace("Matrix", quietly=TRUE)) {
data <- rep(0, 100)
data[(1:10) ^ 2] <- 1:10 * pi
m <- matrix(data, ncol = 20, byrow = TRUE)
m <- Matrix::Matrix(m, sparse = TRUE)
m_hf <- as.h2o(m)
stopifnot(is.h2o(m_hf), dim(m_hf) == dim(m))
}
as.matrix.H2OFrame  

Convert an H2OFrame to a matrix

Description

Convert an H2OFrame to a matrix

Usage

## S3 method for class 'H2OFrame'

as.matrix(x, ...)

Arguments

x  
An H2OFrame object

...  
Further arguments to be passed down from other methods.

Examples

## Not run:
library(h2o)
h2o.init()
iris_hf <- as.h2o(iris)
describe <- h2o.describe(iris_hf)
mins = as.matrix(apply(iris_hf, 2, min))
print(mins)

## End(Not run)

as.numeric  

Convert H2O Data to Numeric

Description

Converts an H2O column into a numeric value column. If the column type is enum and you want to convert it to numeric, you should first convert it to character then convert it to numeric. Otherwise, the values may be converted to underlying factor values, not the expected mapped values.

Usage

as.numeric(x)

Arguments

x  
a column from an H2OFrame data set.
Examples

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
prostate[, 2] <- as.factor (prostate[, 2])
prostate[, 2] <- as.numeric(prostate[, 2])

## End(Not run)
```

---

### as.vector.H2OFrame

Convert an H2OFrame to a vector

#### Description

Convert an H2OFrame to a vector

#### Usage

```
## S3 method for class 'H2OFrame'
as.vector(x, mode)
```

#### Arguments

- `x`  
  - An H2OFrame object

- `mode`  
  - Mode to coerce vector to

#### Examples

```r
## Not run:
library(h2o)
h2o.init()
iris_hf <- as.h2o(iris)
cor_R <- cor(as.matrix(iris[, 1]))
cor_h2o <- cor(iris_hf[, 1])
iris_R_cor <- cor(iris[, 1:4])
iris_H2O_cor <- as.data.frame(cor(iris_hf[, 1:4]))
h2o_vec <- as.vector(unlist(iris_H2O_cor))
r_vec <- as.vector(unlist(iris_R_cor))

## End(Not run)
```
Australia Coastal Data

**Description**
Temperature, soil moisture, runoff, and other environmental measurements from the Australia coast. The data is available from https://cs.colby.edu/courses/S11/cs251/labs/lab07/AustraliaSubset.csv.

**Format**
A data frame with 251 rows and 8 columns

---

**colnames**

*Returns the column names of an H2OFrame*

**Description**
Returns the column names of an H2OFrame

**Usage**

```r
colnames(x, do.NULL = TRUE, prefix = "col")
```

**Arguments**

- `x` An H2OFrame object.
- `do.NULL` logical. If FALSE and names are NULL, names are created.
- `prefix` for created names.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

iris_hf <- as.h2o(iris)
colnames(iris_hf) # Returns "Sepal.Length" "Sepal.Width" "Petal.Length" "Petal.Width" "Species"

## End(Not run)
```
**dim.H2OFrame**

---

**dim.H2OFrame**

*Returns the Dimensions of an H2OFrame*

---

**Description**

Returns the number of rows and columns for an H2OFrame object.

**Usage**

```r
## S3 method for class 'H2OFrame'
dim(x)
```

**Arguments**

- `x`: An H2OFrame object.

**See Also**

- `dim` for the base R method.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

iris_hf <- as.h2o(iris)
dim(iris_hf)
## End(Not run)
```

---

**dimnames.H2OFrame**

*Column names of an H2OFrame*

---

**Description**

Set column names of an H2O Frame.

**Usage**

```r
## S3 method for class 'H2OFrame'
dimnames(x)
```

**Arguments**

- `x`: An H2OFrame
Examples

```r
# Not run:
library(h2o)
h2o.init()

n <- 2000
# Generate variables V1, ... V10
X <- matrix(rnorm(10 * n), n, 10)
# y = +1 if sum_i x_{ij}^2 > chisq median on 10 df
y <- rep(-1, n)
y[apply(X*X, 1, sum) > qchisq(.5, 10)] <- 1
# Assign names to the columns of X:
dimnames(X)[[2]] <- c("V1", "V2", "V3", "V4", "V5", "V6", "V7", "V8", "V9", "V10")

## End(Not run)
```

---

feature_frequencies.H2OModel

Retrieve the number of occurrences of each feature for given observations Available for GBM, Random Forest and Isolation Forest models.

Description

Retrieve the number of occurrences of each feature for given observations Available for GBM, Random Forest and Isolation Forest models.

Usage

```r
feature_frequencies.H2OModel(object, newdata, ...)
h2o.feature_frequencies(object, newdata, ...)
```

Arguments

- `object` a fitted `H2OModel` object for which prediction is desired
- `newdata` An H2OFrame object in which to look for variables with which to predict.
- `...` additional arguments to pass on.

Value

Returns an H2OFrame contain per-feature frequencies on the predict path for each input row.

See Also

- `h2o.gbm` and `h2o.randomForest` for model generation in h2o.
generate_col_ind

CHeck to see if the column names/indices entered is valid for the dataframe given. This is an internal function.

Description

CHeck to see if the column names/indices entered is valid for the dataframe given. This is an internal function.

Usage

generate_col_ind(data, by)

Arguments

data The H2OFrame whose column names or indices are entered as a list.
by The column names/indices in a list.

generate_col_ind

get_seed.H2OModel

Get the seed from H2OModel which was used during training. If a user does not set the seed parameter before training, the seed is autogenerated. It returns seed as the string if the value is bigger than the integer. For example, an autogenerated seed is always long so that the seed in R is a string.

Description

Get the seed from H2OModel which was used during training. If a user does not set the seed parameter before training, the seed is autogenerated. It returns seed as the string if the value is bigger than the integer. For example, an autogenerated seed is always long so that the seed in R is a string.

Usage

generate_col_ind(data, by)

generate_col_ind

get_seed.H2OModel(object)

h2o.get_seed(object)

Arguments

object a fitted H2OModel object.

Value

Returns seed to be used during training a model. Could be numeric or string.
Compute the absolute value of x

Description
Compute the absolute value of x

Usage

h2o.abs(x)

Arguments

x An H2OFrame object.

See Also

MathFun for the base R implementation, abs().

Examples

## Not run:
library(h2o)
h2o.init()
smtrees_hf <- h2o.importFile(url)
model <- h2o.gbm(x = c("girth", "height"), y = "vol", ntrees = 3, max_depth = 1,
    distribution = "gaussian", min_rows = 2, learn_rate = .1,
    training_frame = smtrees_hf)
pred <- as.data.frame(predict(model, newdata = smtrees_hf))
diff <- pred - smtrees_df[, 4]
diff_abs <- abs(diff)
print(diff_abs)
## End(Not run)
h2o.acos  

Compute the arc cosine of x

Description
Compute the arc cosine of x

Usage
h2o.acos(x)

Arguments
x  An H2OFrame object.

See Also
Trig for the base R implementation, acos().

Examples
## Not run:
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
h2o.acos(prostate[, 2])
## End(Not run)

h2o.aggregated_frame  

Retrieve an aggregated frame from an Aggregator model

Description
Retrieve an aggregated frame from the Aggregator model and use it to create a new frame.

Usage
h2o.aggregated_frame(model)

Arguments
model  an H2OClusteringModel corresponding from a h2o.aggregator call.
Examples

## Not run:
library(h2o)
h2o.init()
df <- h2o.createFrame(rows = 100,
cols = 5,
categorical_fraction = 0.6,
integer_fraction = 0,
binary_fraction = 0,
real_range = 100,
integer_range = 100,
missing_fraction = 0)
target_num_exemplars = 1000
rel_tol_num_exemplars = 0.5
encoding = "Eigen"
agg <- h2o.aggregator(training_frame = df,
target_num_exemplars = target_num_exemplars,
rel_tol_num_exemplars = rel_tol_num_exemplars,
categorical_encoding = encoding)
# Use the aggregated frame to create a new dataframe
new_df <- h2o.aggregated_frame(agg)

## End(Not run)

---

**h2o.aggregator**  
Build an Aggregated Frame

**Description**

Builds an Aggregated Frame of an H2OFrame.

**Usage**

```r
h2o.aggregator(
  training_frame,
  x,
  model_id = NULL,
  ignore_const_cols = TRUE,
  target_num_exemplars = 5000,
  rel_tol_num_exemplars = 0.5,
  transform = c("NONE", "STANDARDIZE", "NORMALIZE", "DEMEAN", "DESCALE"),
  categorical_encoding = c("AUTO", "Enum", "OneHotInternal", "OneHotExplicit",
                           "Binary", "Eigen", "LabelEncoder", "SortByResponse", "EnumLimited"),
  save_mapping_frame = FALSE,
  num_iteration_without_new_exemplar = 500,
  export_checkpoints_dir = NULL
)
```
Arguments

- **training_frame**  
  Id of the training data frame.

- **x**  
  A vector containing the character names of the predictors in the model.

- **model_id**  
  Destination id for this model; auto-generated if not specified.

- **ignore_const_cols**  
  Logical. Ignore constant columns. Defaults to TRUE.

- **target_num_exemplars**  
  Targeted number of exemplars. Defaults to 5000.

- **rel_tol_num_exemplars**  
  Relative tolerance for number of exemplars (e.g., 0.5 is +/- 50 percents). Defaults to 0.5.

- **transform**  
  Transformation of training data. Must be one of: "NONE", "STANDARDIZE", "NORMALIZE", "DEMEAN", "DESCALE". Defaults to NORMALIZE.

- **categorical_encoding**  
  Encoding scheme for categorical features. Must be one of: "AUTO", "Enum", "OneHotInternal", "OneHotExplicit", "Binary", "Eigen", "LabelEncoder", "Sort-ByResponse", "EnumLimited". Defaults to AUTO.

- **save_mapping_frame**  
  Logical. Whether to export the mapping of the aggregated frame. Defaults to FALSE.

- **num_iteration_without_new_exemplar**  
  The number of iterations to run before aggregator exits if the number of exemplars collected didn’t change. Defaults to 500.

- **export_checkpoints_dir**  
  Automatically export generated models to this directory.

Examples

```r
## Not run:
library(h2o)
h2o.init()
df <- h2o.createFrame(rows = 100,
cols = 5,
categorical_fraction = 0.6,
integer_fraction = 0,
binary_fraction = 0,
real_range = 100,
integer_range = 100,
missing_fraction = 0)

target_num_exemplars = 1000
rel_tol_num_exemplars = 0.5
encoding = "Eigen"
agg <- h2o.aggregator(training_frame = df,
target_num_exemplars = target_num_exemplars,
rel_tol_num_exemplars = rel_tol_num_exemplars,
categorical_encoding = encoding)

## End(Not run)
```
h2o.aic

Retrieve the Akaike information criterion (AIC) value

Description

Retrieves the AIC value. If "train", "valid", and "xval" parameters are FALSE (default), then the training AIC value is returned. If more than one parameter is set to TRUE, then a named vector of AICs are returned, where the names are "train", "valid" or "xval".

Usage

h2o.aic(object, train = FALSE, valid = FALSE, xval = FALSE)

Arguments

- **object**: An H2OModel or H2OModelMetrics.
- **train**: Retrieve the training AIC
- **valid**: Retrieve the validation AIC
- **xval**: Retrieve the cross-validation AIC

Examples

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
p_sid <- h2o.runif(prostate)
prostate_train <- prostate[p_sid > .2,]
prostate_glm <- h2o.glm(x = 3:7, y = 2, training_frame = prostate_train)
aic_basic <- h2o.aic(prostate_glm)
print(aic_basic)
## End(Not run)
```

h2o.all

Given a set of logical vectors, are all of the values true?

Description

Given a set of logical vectors, are all of the values true?

Usage

h2o.all(x)
**Arguments**

- `object` An H2OAutoEncoderModel object that represents the model to be used for anomaly detection.
- `data` An H2OFrame object.
- `per_feature` Whether to return the per-feature squared reconstruction error

**Value**

Returns an H2OFrame object containing the reconstruction MSE or the per-feature squared error.

**See Also**

- `h2o.deeplearning` for making an H2OAutoEncoderModel.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
prostate_path = system.file("extdata", "prostate.csv", package = "h2o")
prostate = h2o.importFile(path = prostate_path)
prostate_dl = h2o.deeplearning(x = 3:9, training_frame = prostate, autoencoder = TRUE,
                               hidden = c(10, 10), epochs = 5)
prostate_anon = h2o.anomaly(prostate_dl, prostate)
head(prostate_anon)

prostate_anon_per_feature = h2o.anomaly(prostate_dl, prostate, per_feature = TRUE)
head(prostate_anon_per_feature)
## End(Not run)
```
**H2O ANOVAGLM** is used to calculate Type III SS which is used to evaluate the contributions of individual predictors and their interactions to a model. Predictors or interactions with negligible contributions to the model will have high p-values while those with more contributions will have low p-values.

### Description

H2O ANOVAGLM is used to calculate Type III SS which is used to evaluate the contributions of individual predictors and their interactions to a model. Predictors or interactions with negligible contributions to the model will have high p-values while those with more contributions will have low p-values.

### Usage

```r
h2o.anovaglm(
  x,
  y,
  training_frame,
  model_id = NULL,
  seed = -1,
  ignore_const_cols = TRUE,
  score_each_iteration = FALSE,
  offset_column = NULL,
  weights_column = NULL,
  family = c("AUTO", "gaussian", "binomial", "fractionalbinomial", "quasibinomial", "poisson", "gamma", "tweedie", "negativebinomial"),
  tweedie_variance_power = 0,
  tweedie_link_power = 1,
  theta = 0,
  solver = c("AUTO", "IRLSM", "L_BFGS", "COORDINATE_DESCENT_NAIVE",
             "COORDINATE_DESCENT", "GRADIENT_DESCENT_LH", "GRADIENT_DESCENT_SQERR"),
  missing_values_handling = c("MeanImputation", "Skip", "PlugValues"),
  plug_values = NULL,
  compute_p_values = TRUE,
  standardize = TRUE,
  non_negative = FALSE,
  max_iterations = 0,
  link = c("family_default", "identity", "logit", "log", "inverse", "tweedie",
           "ologit"),
  prior = 0,
  alpha = NULL,
  lambda = c(0),
  lambda_search = FALSE,
  stopping_rounds = 0,
  stopping_metric = c("AUTO", "deviance", "logloss", "MSE", "RMSE", "MAE", "RMSLE"),
)```
"AUC", "AUCPR", "lift_top_group", "misclassification", "mean_per_class_error", "custom", "custom_increasing"),
  early_stopping = FALSE,
  stopping_tolerance = 0.001,
  balance_classes = FALSE,
  class_sampling_factors = NULL,
  max_after_balance_size = 5,
  max_runtime_secs = 0,
  save_transformed_framekeys = FALSE,
  highest_interaction_term = 0,
  nparallelism = 4,
  type = 0
)

Arguments

x  (Optional) A vector containing the names or indices of the predictor variables to use in building the model. If x is missing, then all columns except y are used.

y  The name or column index of the response variable in the data. The response must be either a numeric or a categorical/factor variable. If the response is numeric, then a regression model will be trained, otherwise it will train a classification model.

training_frame  Id of the training data frame.

model_id  Destination id for this model; auto-generated if not specified.

seed  Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).

ignore_const_cols  Logical. Ignore constant columns. Defaults to TRUE.

score_each_iteration  Logical. Whether to score during each iteration of model training. Defaults to FALSE.

offset_column  Offset column. This will be added to the combination of columns before applying the link function.

weights_column  Column with observation weights. Giving some observation a weight of zero is equivalent to excluding it from the dataset; giving an observation a relative weight of 2 is equivalent to repeating that row twice. Negative weights are not allowed. Note: Weights are per-row observation weights and do not increase the size of the data frame. This is typically the number of times a row is repeated, but non-integer values are supported as well. During training, rows with higher weights matter more, due to the larger loss function pre-factor. If you set weight = 0 for a row, the returned prediction frame at that row is zero and this is incorrect. To get an accurate prediction, remove all rows with weight == 0.

family  Family. Use binomial for classification with logistic regression, others are for regression problems. Must be one of: "AUTO", "gaussian", "binomial", "fractionalbinomial", "quasibinomial", "poisson", "gamma", "tweedie", "negativebinomial". Defaults to AUTO.
tweedie_variance_power
  Tweedie variance power Defaults to 0.

tweedie_link_power
  Tweedie link power Defaults to 1.

theta
  Theta Defaults to 0.

solver
  AUTO will set the solver based on given data and the other parameters. IRLSM
  is fast on problems with small number of predictors and for lambda-search
  with L1 penalty, L_BFGS scales better for datasets with many columns. Must be
  one of: "AUTO", "IRLSM", "L_BFGS", "COORDINATE_DESCENT_NAIVE",
  "COORDINATE_DESCENT", "GRADIENT_DESCENT_LH", "GRADIENT_DESCENT_SQERR".
  Defaults to IRLSM.

missing_values_handling
  Handling of missing values. Either MeanImputation, Skip or PlugValues. Must
  be one of: "MeanImputation", "Skip", "PlugValues". Defaults to MeanImputation.

plug_values
  Plug Values (a single row frame containing values that will be used to im-
  pute missing values of the training/validation frame, use with conjunction miss-
  ing_values_handling = PlugValues)

compute_p_values
  Logical. Request p-values computation, p-values work only with IRLSM solver
  and no regularization Defaults to TRUE.

standardize
  Logical. Standardize numeric columns to have zero mean and unit variance
  Defaults to TRUE.

non_negative
  Logical. Restrict coefficients (not intercept) to be non-negative Defaults to
  FALSE.

max_iterations
  Maximum number of iterations Defaults to 0.

link
  Link function. Must be one of: "family_default", "identity", "logit", "log",
  "inverse", "tweedie", "ologit". Defaults to family_default.

prior
  Prior probability for y==1. To be used only for logistic regression iff the data
  has been sampled and the mean of response does not reflect reality. Defaults to
  0.

alpha
  Distribution of regularization between the L1 (Lasso) and L2 (Ridge) penalties.
  A value of 1 for alpha represents Lasso regression, a value of 0 produces Ridge
  regression, and anything in between specifies the amount of mixing between the
  two. Default value of alpha is 0 when SOLVER = 'L-BFGS'; 0.5 otherwise.

lambda
  Regularization strength Defaults to c(0.0).

lambda_search
  Logical. Use lambda search starting at lambda max, given lambda is then in-
  terpreted as lambda min Defaults to FALSE.

stopping_rounds
  Early stopping based on convergence of stopping_metric. Stop if simple moving
  average of length k of the stopping_metric does not improve for k:=stopping_rounds
  scoring events (0 to disable) Defaults to 0.

stopping_metric
  Metric to use for early stopping (AUTO: logloss for classification, deviance
  for regression and anomoly_score for Isolation Forest). Note that custom
and custom_increasing can only be used in GBM and DRF with the Python client. Must be one of: "AUTO", "deviance", "logloss", "MSE", "RMSE", "MAE", "RMSLE", "AUC", "AUCPR", "lift_top_group", "misclassification", "mean_per_class_error", "custom", "custom_increasing". Defaults to AUTO.

**early_stopping**

Logical. Stop early when there is no more relative improvement on train or validation (if provided). Defaults to FALSE.

**stopping_tolerance**

Relative tolerance for metric-based stopping criterion (stop if relative improvement is not at least this much) Defaults to 0.001.

**balance_classes**

Logical. Balance training data class counts via over/under-sampling (for imbalanced data). Defaults to FALSE.

**class_sampling_factors**

Desired over/under-sampling ratios per class (in lexicographic order). If not specified, sampling factors will be automatically computed to obtain class balance during training. Requires balance_classes.

**max_after_balance_size**

Maximum relative size of the training data after balancing class counts (can be less than 1.0). Requires balance_classes. Defaults to 5.0.

**max_runtime_secs**

Maximum allowed runtime in seconds for model training. Use 0 to disable. Defaults to 0.

**save_transformed_framekeys**

Logical. true to save the keys of transformed predictors and interaction column. Defaults to FALSE.

**highest_interaction_term**

Limit the number of interaction terms, if 2 means interaction between 2 columns only, 3 for three columns and so on... Default to 2. Defaults to 0.

**nparallelism**

Number of models to build in parallel. Default to 4. Adjust according to your system. Defaults to 4.

**type**

Refer to the SS type 1, 2, 3, or 4. We are currently only supporting 3 Defaults to 0.

---

**Examples**

```r
## Not run:
h2o.init()

# Run ANOVA GLM of VOL ~ CAPSULE + RACE
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
prostate$CAPSULE <- as.factor(prostate$CAPSULE)
model <- h2o.anovaglm(y = "VOL", x = c("CAPSULE","RACE"), training_frame = prostate)

## End(Not run)
```
h2o.any

Given a set of logical vectors, is at least one of the values true?

Description

Given a set of logical vectors, is at least one of the values true?

Usage

h2o.any(x)

Arguments

x An H2OFrame object.

See Also

all for the base R implementation.

Examples

## Not run:
library(h2o)
h2o.init()

iris <- h2o.importFile(f)
h2o.any(iris[, 1] < 1000)

## End(Not run)

h2o.anyFactor

Check H2OFrame columns for factors

Description

Determines if any column of an H2OFrame object contains categorical data.

Usage

h2o.anyFactor(x)

Arguments

x An H2OFrame object.
Value

Returns a logical value indicating whether any of the columns in x are factors.

Examples

```r
## Not run:
library(h2o)
h2o.init()
iris_hf <- as.h2o(iris)
h2o.anyFactor(iris_hf)

## End(Not run)
```

---

**h2o.api**

*Perform a REST API request to a previously connected server.*

---

Description

This function is mostly for internal purposes, but may occasionally be useful for direct access to the backend H2O server. It has same parameters as :meth:`H2OConnection.request <h2o.backend.H2OConnection.request>`.

Usage

```r
h2o.api(endpoint, params = NULL, json = NULL)
```

Arguments

- **endpoint**: A H2O REST API endpoint.
- **params**: A list of params passed in the url.
- **json**: A list of params passed as a json payload.

Details

REST API endpoints can be obtained using:

```r
endpoints <- sapply(h2o.api("GET /3/Metadata/endpoints")$routes, function(r) paste(r$http_method, r$url_pattern))
```

For a given route, the supported params can be obtained using:

```r
parameters <- sapply(h2o.api("GET /3/Metadata/schemas/{route$input_schema}")$schemas[[1]]$fields, function(f) { l <- list(); l[f$name] <- f$help; l })
```

Value

The parsed response.
h2o.arrange

Sorts an H2O frame by columns

Examples

```r
## Not run:
res <- h2o.api("GET /3/NetworkTest")
res$table

## End(Not run)
```

## Description

Sorts H2OFrame by the columns specified. H2OFrame can contain String columns but should not sort on any String columns. Otherwise, an error will be thrown. To sort column c1 in descending order, do desc(c1). Returns a new H2OFrame, like dplyr::arrange.

## Usage

```r
h2o.arrange(x, ...)
```

## Arguments

- **x**: The H2OFrame input to be sorted.
- **...**: The column names to sort by.

## Examples

```r
## Not run:
library(h2o)
h2o.init()

iris <- h2o.importFile(f)
h2o.arrange(iris, "species","petal_len","petal_wid")

## End(Not run)
```
h2o.ascharacter

Convert H2O Data to Characters

Description
Convert H2O Data to Characters

Usage
h2o.ascharacter(x)

Arguments
x An H2OFrame object.

Examples
## Not run:
library(h2o)
h2o.init()
iris <- h2o.importFile(f)
h2o.ascharacter(iris["species"])
## End(Not run)

h2o.asfactor

Convert H2O Data to Factors

Description
Convert H2O Data to Factors

Usage
h2o.asfactor(x)

Arguments
x An H2OFrame object.

See Also
factor for the base R implementation, as.factor().
Examples

```r
## Not run:
library(h2o)
h2o.init()

cars <- h2o.importFile(f)
h2o.asfactor(cars[cylinders])

## End(Not run)
```

### h2o.asnumeric

**Convert H2O Data to Numerics**

**Description**

If the column type is enum and you want to convert it to numeric, you should first convert it to character then convert it to numeric. Otherwise, the values may be converted to underlying factor values, not the expected mapped values.

**Usage**

```r
h2o.asnumeric(x)
```

**Arguments**

- `x` An H2OFrame object.

**See Also**

`numeric` for the base R implementation, `as.numeric()`.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

cars <- h2o.importFile(f)
h2o.ascharacter(cars)
h2o.asnumeric(cars)

## End(Not run)
```
h2o.assign  Rename an H2O object.

Description

Makes a copy of the data frame and gives it the desired key.

Usage

h2o.assign(data, key)

Arguments

data An H2OFrame object
key The key to be associated with the H2O parsed data object

Examples

## Not run:
library(h2o)
h2o.init()

cars <- h2o.importFile(f)
dim(cars)
split <- h2o.splitFrame(data = cars, ratios = 0.8)
train <- h2o.assign(split[[1]], key = "train")
test <- h2o.assign(split[[2]], key = "test")
dim(train)
dim(test)

## End(Not run)

h2o.as_date  Convert between character representations and objects of Date class

Description

Functions to convert between character representations and objects of class "Date" representing calendar dates.

Usage

h2o.as_date(x, format, ...)
Arguments

x    H2OFrame column of strings or factors to be converted
format A character string indicating date pattern
...  Further arguments to be passed from or to other methods.

h2o.auc

Retrieve the AUC

Description

Retrieves the AUC value from an H2OBinomialMetrics. If "train", "valid", and "xval" parameters are FALSE (default), then the training AUC value is returned. If more than one parameter is set to TRUE, then a named vector of AUCs are returned, where the names are "train", "valid" or "xval".

Usage

h2o.auc(object, train = FALSE, valid = FALSE, xval = FALSE)

Arguments

object    An H2OBinomialMetrics or H2OMultinomialMetrics object.
train     Retrieve the training AUC
valid     Retrieve the validation AUC
xval      Retrieve the cross-validation AUC

See Also

h2o.giniCoef for the Gini coefficient, h2o.mse for MSE, and h2o.metric for the various threshold metrics. See h2o.performance for creating H2OModelMetrics objects.

Examples

## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(prostate_path)

prostate[, 2] <- as.factor(prostate[, 2])
model <- h2o.gbm(x = 3:9, y = 2, training_frame = prostate, distribution = "bernoulli")
perf <- h2o.performance(model, prostate)
h2o.auc(perf)

## End(Not run)
Description

Retrieves the AUCPR value from an H2OBinomialMetrics. If "train", "valid", and "xval" parameters are FALSE (default), then the training AUCPR value is returned. If more than one parameter is set to TRUE, then a named vector of AUCPRs are returned, where the names are "train", "valid" or "xval".

Usage

h2o.aucpr(object, train = FALSE, valid = FALSE, xval = FALSE)

h2o.pr_auc(object, train = FALSE, valid = FALSE, xval = FALSE)

Arguments

object  
An H2OBinomialMetrics object.

train  
Retrieve the training aucpr

valid  
Retrieve the validation aucpr

xval  
Retrieve the cross-validation aucpr

See Also

h2o.giniCoef for the Gini coefficient, h2o.mse for MSE, and h2o.metric for the various threshold metrics. See h2o.performance for creating H2OModelMetrics objects.

Examples

## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(prostate_path)

prostate[, 2] <- as.factor(prostate[, 2])
model <- h2o.gbm(x = 3:9, y = 2, training_frame = prostate, distribution = "bernoulli")
perf <- h2o.performance(model, prostate)
h2o.aucpr(perf)

## End(Not run)
Description

The Automatic Machine Learning (AutoML) function automates the supervised machine learning model training process. AutoML finds the best model, given a training frame and response, and returns an H2OAutoML object, which contains a leaderboard of all the models that were trained in the process, ranked by a default model performance metric.

Usage

```r
h2o.automl(
  x,
  y,
  training_frame,
  validation_frame = NULL,
  leaderboard_frame = NULL,
  blending_frame = NULL,
  nfolds = -1,
  fold_column = NULL,
  weights_column = NULL,
  balance_classes = FALSE,
  class_sampling_factors = NULL,
  max_after_balance_size = 5,
  max_runtime_secs = NULL,
  max_runtime_secs_per_model = NULL,
  max_models = NULL,
  stopping_metric = c("AUTO", "deviance", "logloss", "MSE", "RMSE", "MAE", "RMSLE", "AUC", "AUCPR", "lift_top_group", "misclassification", "mean_per_class_error"),
  stopping_tolerance = NULL,
  stopping_rounds = 3,
  seed = NULL,
  project_name = NULL,
  exclude_algos = NULL,
  include_algos = NULL,
  modeling_plan = NULL,
  preprocessing = NULL,
  exploitation_ratio = -1,
  monotone_constraints = NULL,
  keep_cross_validation_predictions = FALSE,
  keep_cross_validation_models = FALSE,
  keep_cross_validation_fold_assignment = FALSE,
  sort_metric = c("AUTO", "deviance", "logloss", "MSE", "RMSE", "MAE", "RMSLE", "AUC", "AUCPR", "mean_per_class_error"),
  export_checkpoints_dir = NULL,
  verbosity = "warn",
)```
Arguments

x  A vector containing the names or indices of the predictor variables to use in building the model. If x is missing, then all columns except y are used.

y  The name or index of the response variable in the model. For classification, the y column must be a factor, otherwise regression will be performed. Indexes are 1-based in R.

training_frame  Training frame (H2OFrame or ID).

validation_frame  Validation frame (H2OFrame or ID); Optional. This argument is ignored unless the user sets nfolds = 0. If cross-validation is turned off, then a validation frame can be specified and used for early stopping of individual models and early stopping of the grid searches. By default and when nfolds > 1, cross-validation metrics will be used for early stopping and thus validation_frame will be ignored.

leaderboard_frame  Leaderboard frame (H2OFrame or ID); Optional. If provided, the Leaderboard will be scored using this data frame instead of using cross-validation metrics, which is the default.

blending_frame  Blending frame (H2OFrame or ID) used to train the the metalearning algorithm in Stacked Ensembles (instead of relying on cross-validated predicted values); Optional. When provided, it also is recommended to disable cross validation by setting nfolds=0 and to provide a leaderboard frame for scoring purposes.

nfolds  Number of folds for k-fold cross-validation. Must be >= 2; defaults to 5. Use 0 to disable cross-validation; this will also disable Stacked Ensemble (thus decreasing the overall model performance).

fold_column  Column with cross-validation fold index assignment per observation; used to override the default, randomized, 5-fold cross-validation scheme for individual models in the AutoML run.

weights_column  Column with observation weights. Giving some observation a weight of zero is equivalent to excluding it from the dataset; giving an observation a relative weight of 2 is equivalent to repeating that row twice. Negative weights are not allowed.

balance_classes  Logical. Specify whether to oversample the minority classes to balance the class distribution; only applicable to classification. If the oversampled size of the dataset exceeds the maximum size calculated during max_after_balance_size parameter, then the majority class will be undersampled to satisfy the size limit. Defaults to FALSE.

class_sampling_factors  Desired over/under-sampling ratios per class (in lexicographic order). If not specified, sampling factors will be automatically computed to obtain class balance during training. Requires balance_classes.
max_after_balance_size

- Maximum relative size of the training data after balancing class counts (can be less than 1.0). Requires balance_classes. Defaults to 5.0.

max_runtime_secs

- This argument specifies the maximum time that the AutoML process will run for. If neither max_runtime_secs nor max_models are specified by the user, then max_runtime_secs defaults to 3600 seconds (1 hour).

max_runtime_secs_per_model

- Maximum runtime in seconds dedicated to each individual model training process. Use 0 to disable. Defaults to 0.

max_models

- Maximum number of models to build in the AutoML process (does not include Stacked Ensembles). Defaults to NULL (no strict limit).

stopping_metric

- Metric to use for early stopping ("AUTO" is logloss for classification, deviance for regression). Must be one of "AUTO", "deviance", "logloss", "MSE", "RMSE", "MAE", "RMSLE", "AUC", "AUCPR", "lift_top_group", "misclassification", "mean_per_class_error". Defaults to "AUTO".

stopping_tolerance

- Relative tolerance for metric-based stopping criterion (stop if relative improvement is not at least this much). This value defaults to 0.001 if the dataset is at least 1 million rows; otherwise it defaults to a bigger value determined by the size of the dataset and the non-NA-rate. In that case, the value is computed as 1/sqrt(nrows * non-NA-rate).

stopping_rounds

- Integer. Early stopping based on convergence of stopping_metric. Stop if simple moving average of length k of the stopping_metric does not improve for k (stopping_rounds) scoring events. Defaults to 3 and must be an non-zero integer. Use 0 to disable early stopping.

seed

- Integer. Set a seed for reproducibility. AutoML can only guarantee reproducibility if max_models or early stopping is used because max_runtime_secs is resource limited, meaning that if the resources are not the same between runs, AutoML may be able to train more models on one run vs another. In addition, H2O Deep Learning models are not reproducible by default for performance reasons, so if the user requires reproducibility, then exclude_algos must contain "DeepLearning".

project_name

- Character string to identify an AutoML project. Defaults to NULL, which means a project name will be auto-generated. More models can be trained and added to an existing AutoML project by specifying the same project name in multiple calls to the AutoML function (as long as the same training frame is used in subsequent runs).

exclude_algos

- Vector of character strings naming the algorithms to skip during the model-building phase. An example use is exclude_algos = c("GLM","DeepLearning","DRF"), and the full list of options is: "DRF" (Random Forest and Extremely-Randomized Trees), "GLM", "XGBoost", "GBM", "DeepLearning" and "StackedEnsemble". Defaults to NULL, which means that all appropriate H2O algorithms will be used, if the search stopping criteria allow. Optional.
include_algos Vector of character strings naming the algorithms to restrict to during the model-building phase. This can’t be used in combination with exclude_algos param. Defaults to NULL, which means that all appropriate H2O algorithms will be used, if the search stopping criteria allow. Optional.

modeling_plan List. The list of modeling steps to be used by the AutoML engine (they may not all get executed, depending on other constraints). Optional (Expert usage only).

preprocessing List. The list of preprocessing steps to run. Only 'target_encoding' is currently supported.

exploitation_ratio The budget ratio (between 0 and 1) dedicated to the exploitation (vs exploration) phase. By default, this is set to AUTO (exploitation_ratio=-1) as this is still experimental; to activate it, it is recommended to try a ratio around 0.1. Note that the current exploitation phase only tries to fine-tune the best XGBoost and the best GBM found during exploration.

monotone_constraints List. A mapping representing monotonic constraints. Use +1 to enforce an increasing constraint and -1 to specify a decreasing constraint.

keep_cross_validation_predictions Logical. Whether to keep the predictions of the cross-validation predictions. This needs to be set to TRUE if running the same AutoML object for repeated runs because CV predictions are required to build additional Stacked Ensemble models in AutoML. This option defaults to FALSE.

keep_cross_validation_models Logical. Whether to keep the cross-validated models. Keeping cross-validation models may consume significantly more memory in the H2O cluster. This option defaults to FALSE.

keep_cross_validation_fold_assignment Logical. Whether to keep fold assignments in the models. Deleting them will save memory in the H2O cluster. Defaults to FALSE.

sort_metric Metric to sort the leaderboard by. For binomial classification choose between "AUC", "AUCPR", "logloss", "mean_per_class_error", "RMSE", "MSE". For regression choose between "mean_residual_deviance", "RMSE", "MSE", "MAE", and "RMSLE". For multinomial classification choose between "mean_per_class_error", "logloss", "RMSE", "MSE". Default is "AUTO". If set to "AUTO", then "AUC" will be used for binomial classification, "mean_per_class_error" for multinomial classification, and "mean_residual_deviance" for regression.

export_checkpoints_dir (Optional) Path to a directory where every model will be stored in binary form.

verbosity Verbosity of the backend messages printed during training; Optional. Must be one of NULL (live log disabled), "debug", "info", "warn", "error". Defaults to "warn".

Details

AutoML trains several models, cross-validated by default, by using the following available algorithms:
- XGBoost
- GBM (Gradient Boosting Machine)
- GLM (Generalized Linear Model)
- DRF (Distributed Random Forest)
- XRT (eXtremely Randomized Trees)
- DeepLearning (Fully Connected Deep Neural Network)

It also applies HPO on the following algorithms:

- XGBoost
- GBM
- DeepLearning

In some cases, there will not be enough time to complete all the algorithms, so some may be missing from the leaderboard.

Finally, AutoML also trains several Stacked Ensemble models at various stages during the run. Mainly two kinds of Stacked Ensemble models are trained:

- one of all available models at time t.
- one of only the best models of each kind at time t.

Note that Stacked Ensemble models are trained only if there isn’t another stacked ensemble with the same base models.

Value

An H2OAutoML object.

Examples

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.importFile(path = prostate_path, header = TRUE)
y <- "CAPSULE"
prostate[,y] <- as.factor(prostate[,y])  # convert to factor for classification
aml <- h2o.automl(y = y, training_frame = prostate, max_runtime_secs = 30)
lb <- h2o.get_leaderboard(aml)
head(lb)
```

## End(Not run)
h2o.auuc

Retrieve the default AUUC

Description

Retrieves the AUUC value from an H2OBinomialUpliftMetrics. The type of AUUC depends on auuc_type which was set before training. If you need specific AUUC, see h2o.auuc_table function. If "train" and "valid" parameters are FALSE (default), then the training AUUC value is returned. If more than one parameter is set to TRUE, then a named vector of AUUCs are returned, where the names are "train", "valid".

Usage

h2o.auuc(object, train = FALSE, valid = FALSE)

Arguments

object An H2OBinomialUpliftMetrics
train Retrieve the training AUUC
valid Retrieve the validation AUUC

Examples

## Not run:
library(h2o)
h2o.init()
f <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/uplift/criteo_uplift_13k.csv"
train <- h2o.importFile(f)
train$treatment <- as.factor(train$treatment)
train$conversion <- as.factor(train$conversion)

model <- h2o.upliftRandomForest(training_frame=train, x=sprintf("f%s",seq(0:10)), y="conversion",
                                 ntrees=10, max_depth=5, treatment_column="treatment",
                                 auuc_type="AUTO")

perf <- h2o.performance(model, train=TRUE)
h2o.auuc(perf)

## End(Not run)

h2o.auuc_table

Retrieve the all types of AUUC in a table

Description

Retrieves the all types of AUUC in a table from an H2OBinomialUpliftMetrics. If "train" and "valid" parameters are FALSE (default), then the training AUUC values is returned. If more than one parameter is set to TRUE, then a named vector of AUUCs are returned, where the names are "train", "valid".
Usage

h2o.auuc_table(object, train = FALSE, valid = FALSE)

Arguments

object An H2OBinomialUpliftMetrics

train Retrieve the training AUUC table

valid Retrieve the validation AUUC table

Examples

## Not run:
library(h2o)
h2o.init()
f <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/uplift/criteo_uplift_13k.csv"
train <- h2o.importFile(f)
train$treatment <- as.factor(train$treatment)
train$conversion <- as.factor(train$conversion)

model <- h2o.upliftRandomForest(training_frame=train, x=sprintf("f%s",seq(0:10)), y="conversion",
                                 ntrees=10, max_depth=5, treatment_column="treatment",
                                 auuc_type="AUTO")

perf <- h2o.performance(model, train=TRUE)
h2o.auuc_table(perf)

## End(Not run)

h2o.betweenss

Get the between cluster sum of squares

Description

Get the between cluster sum of squares. If "train", "valid", and "xval" parameters are FALSE (default), then the training betweenss value is returned. If more than one parameter is set to TRUE, then a named vector of betweenss' are returned, where the names are "train", "valid" or "xval".

Usage

h2o.betweenss(object, train = FALSE, valid = FALSE, xval = FALSE)

Arguments

object An H2OClusteringModel object.

train Retrieve the training between cluster sum of squares

valid Retrieve the validation between cluster sum of squares

xval Retrieve the cross-validation between cluster sum of squares
h2o.biases

Examples

## Not run:
library(h2o)
h2o.init()
predictors <- c("sepal.len", "sepal.wid", "petal.len", "petal.wid")
km <- h2o.kmeans(x = predictors, training_frame = fr, k = 3, nfolds = 3)
h2o.betweenss(km, train = TRUE)

## End(Not run)

h2o.biases

Return the respective bias vector

Description

Return the respective bias vector

Usage

h2o.biases(object, vector_id = 1)

Arguments

object An H2OModel or H2OModelMetrics
vector_id An integer, ranging from 1 to number of layers + 1, that specifies the bias vector to return.

Examples

## Not run:
library(h2o)
h2o.init()

f <- "https://h2o-public-test-data.s3.amazonaws.com/smalldata/chicago/chicagoCensus.csv"
census <- h2o.importFile(f)
census[, 1] <- as.factor(census[, 1])
dl_model <- h2o.deeplearning(x = c(1:3), y = 4, training_frame = census,
hidden = c(17, 191),
epochs = 1,
balance_classes = FALSE,
export_weights_and_biases = TRUE)
h2o.biases(dl_model, vector_id = 1)

## End(Not run)
h2o.bottomN  

**Description**

bottomN function will will grab the bottom N percent of values of a column and return it in a H2OFrame. Extract the top N percent of values of a column and return it in a H2OFrame.

**Usage**

\[ \text{h2o.bottomN}(x, \text{column}, \text{nPercent}) \]

**Arguments**

- \( x \) an H2OFrame
- \( \text{column} \) is a column name or column index to grab the top N percent value from
- \( \text{nPercent} \) is a bottom percentage value to grab

**Value**

An H2OFrame with 2 columns. The first column is the original row indices, second column contains the bottomN values

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

f2 <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/jira/Bottom20Per.csv.zip"
data_Frame <- h2o.importFile(f1)
bottom_Answer <- h2o.importFile(f2)
nPercent <- c(1, 2, 3, 4)
frame_Names <- names(data_Frame)
nP <- nPercent[sample(1:length(nPercent), 1, replace = FALSE)]
col_Index <- sample(1:length(frame_Names), 1, replace = FALSE)
h2o.bottomN(data_Frame, frame_Names[col_Index], nP)

## End(Not run)
```
**h2o.cbind**  
*Combine H2O Datasets by Columns*

**Description**

Takes a sequence of H2O data sets and combines them by column.

**Usage**

```r
h2o.cbind(...)```

**Arguments**

```r
...```

A sequence of H2OFrame arguments. All datasets must exist on the same H2O instance (IP and port) and contain the same number of rows.

**Value**

An H2OFrame object containing the combined...arguments column-wise.

**See Also**

`cbind` for the base R method.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
prostate_cbind <- h2o.cbind(prostate, prostate)
head(prostate_cbind)
## End(Not run)
```

---

**h2o.ceiling**  
*Take a single numeric argument and return a numeric vector with the smallest integers*

**Description**

ceiling takes a single numeric argument x and returns a numeric vector containing the smallest integers not less than the corresponding elements of x.
### h2o.centers

**Usage**

\[ \text{h2o.centers}(\text{object}) \]

**Arguments**

- **object**: An \texttt{H2OClusteringModel} object.

**Description**

Retrieve the Model Centers

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
h2o.centers(fr)
## End(Not run)
```
h2o.centersSTD

**Retrieve the Model Centers STD**

**Description**

Retrieve the Model Centers STD

**Usage**

h2o.centersSTD(object)

**Arguments**

object  
An H2OClusteringModel object.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
predictors <- c("sepal_len", "sepal_wid", "petal_len", "petal_wid")
km <- h2o.kmeans(x = predictors, training_frame = fr, k = 3, nfolds = 3)
h2o.centersSTD(km)
## End(Not run)
```

h2o.centroid_stats

**Retrieve centroid statistics**

**Description**

Retrieve the centroid statistics. If "train" and "valid" parameters are FALSE (default), then the training centroid stats value is returned. If more than one parameter is set to TRUE, then a named list of centroid stats data frames are returned, where the names are "train" or "valid" For cross validation metrics this statistics are not available.

**Usage**

h2o.centroid_stats(object, train = FALSE, valid = FALSE)

**Arguments**

object  
An H2OClusteringModel object.

train  
Retrieve the training centroid statistics

valid  
Retrieve the validation centroid statistics
h2o.clearLog

Delete All H2O R Logs

Description

Clear all H2O R command and error response logs from the local disk. Used primarily for debugging purposes.

Usage

h2o.clearLog()

See Also

h2o.startLogging, h2o.stopLogging, h2o.openLog

Examples

## Not run:
library(h2o)
h2o.init()
predictors <- c("sepal_len", "sepal_wid", "petal_len", "petal_wid")
km <- h2o.kmeans(x = predictors, training_frame = fr, k = 3, nfolds = 3)
h2o.centroid_stats(km, train = TRUE)

## End(Not run)

## Not run:
library(h2o)
australia_path = system.file("extdata", "australia.csv", package = "h2o")
australia = h2o.importFile(path = australia_path)
h2o.stopLogging()
h2o.clearLog()

## End(Not run)
h2o.clusterInfo  
Print H2O cluster info

Description
Print H2O cluster info

Usage
h2o.clusterInfo()

h2o.clusterIsUp  
Determine if an H2O cluster is up or not

Description
Determine if an H2O cluster is up or not

Usage
h2o.clusterIsUp(conn = h2o.getConnection())

Arguments
conn  
H2OConnection object

Value
TRUE if the cluster is up; FALSE otherwise

h2o.clusterStatus  
Return the status of the cluster

Description
Retrieve information on the status of the cluster running H2O.

Usage
h2o.clusterStatus()

See Also
H2OConnection, h2o.init
Examples

```r
## Not run:
h2o.init()
h2o.clusterStatus()

## End(Not run)
```

### h2o.cluster_sizes

**Retrieve the cluster sizes**

#### Description

Retrieve the cluster sizes. If "train", "valid", and "xval" parameters are FALSE (default), then the training cluster sizes value is returned. If more than one parameter is set to TRUE, then a named list of cluster size vectors are returned, where the names are "train", "valid" or "xval".

#### Usage

```r
h2o.cluster_sizes(object, train = FALSE, valid = FALSE, xval = FALSE)
```

#### Arguments

- `object`: An H2OClusteringModel object.
- `train`: Retrieve the training cluster sizes
- `valid`: Retrieve the validation cluster sizes
- `xval`: Retrieve the cross-validation cluster sizes

#### Examples

```r
## Not run:
library(h2o)
h2o.init()
predictors <- c("sepal_len", "sepal_wid", "petal_len", "petal_wid")
km <- h2o.kmeans(x = predictors, training_frame = fr, k = 3, nfolds = 3)
h2o.cluster_sizes(km, train = TRUE)

## End(Not run)
```
Return the coefficients that can be applied to the non-standardized data.

Description

Note: standardize = True by default. If set to False, then coef() returns the coefficients that are fit directly.

Usage

h2o.coef(object, predictorSize = -1)

Arguments

object an H2OModel object.
predictorSize predictor subset size. If specified, will only return model coefficients of that subset size. If not specified will return a lists of model coefficient dicts for all predictor subset size.

Examples

## Not run:
library(h2o)
h2o.init()
cars <- h2o.importFile(f)
predictors <- c("displacement", "power", "weight", "acceleration", "year")
response <- "cylinders"
cars_split <- h2o.splitFrame(data = cars, ratios = 0.8, seed = 1234)
train <- cars_split[[1]]
valid <- cars_split[[2]]
cars_glm <- h2o.glm(balance_classes = TRUE,
seed = 1234,
x = predictors,
y = response,
training_frame = train,
validation_frame = valid)
h2o.coef(cars_glm)

## End(Not run)
h2o.coef_norm

Return coefficients fitted on the standardized data (requires standardize = True, which is on by default). These coefficients can be used to evaluate variable importance.

Description

Return coefficients fitted on the standardized data (requires standardize = True, which is on by default). These coefficients can be used to evaluate variable importance.

Usage

h2o.coef_norm(object, predictorSize = -1)

Arguments

object
   an H2OModel object.
predictorSize
   predictor subset size. If specified, will only return model coefficients of that subset size. If not specified will return a lists of model coefficient dicts for all predictor subset size.

Examples

```r
## Not run:
library(h2o)
h2o.init()

cars <- h2o.importFile(f)
predictors <- c("displacement", "power", "weight", "acceleration", "year")
response <- "cylinders"
cars_split <- h2o.splitFrame(data = cars, ratios = 0.8, seed = 1234)
train <- cars_split[[1]]
valid <- cars_split[[2]]
cars glm <- h2o.glm(balance_classes = TRUE,
   seed = 1234,
   x = predictors,
   y = response,
   training_frame = train,
   validation_frame = valid)
h2o.coef_norm(cars glm)

## End(Not run)
```
**h2o.colnames**

Return column names of an H2OFrame

**Description**

Return column names of an H2OFrame

**Usage**

```r
def h2o.colnames(x)
```

**Arguments**

- `x` An H2OFrame object.

**See Also**

`colnames` for the base R implementation.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)
h2o.colnames(frame)
## End(Not run)
```

**h2o.columns_by_type**

Obtain a list of columns that are specified by `coltype`

**Description**

Obtain a list of columns that are specified by `coltype`

**Usage**

```r
h2o.columns_by_type(object, coltype = "numeric", ...)
```
Arguments

object : H2OFrame object

coltype : A character string indicating which column type to filter by. This must be one of the following: "numeric" - Numeric, but not categorical or time "categorical" - Integer, with a categorical/factor String mapping "string" - String column "time" - Long msec since the Unix Epoch - with a variety of display/parse options "uuid" - UUID "bad" - No none-NA rows (triple negative! all NAs or zero rows)

... Ignored

Value

A list of column indices that correspond to "type"

Examples

```r
## Not run:
# h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
h2o.columns_by_type(prostate, coltype = "numeric")

## End(Not run)
```

h2o.computeGram

*Compute weighted gram matrix.*

Description

Compute weighted gram matrix.

Usage

```r
h2o.computeGram(
  X,
  weights = ",",
  use_all_factor_levels = FALSE,
  standardize = TRUE,
  skip_missing = FALSE
)
```

Arguments

X : an H2OModel corresponding to H2O framel.

weights : character corresponding to name of weight vector in frame.

use_all_factor_levels : logical flag telling h2o whether or not to skip first level of categorical variables during one-hot encoding.
h2o.confusionMatrix

standardize  logical flag telling h2o whether or not to standardize data
skip_missing logical flag telling h2o whether skip rows with missing data or impute them with mean

---

**Description**

Retrieve either a single or many confusion matrices from H2O objects.

**Usage**

```r
h2o.confusionMatrix(object, ...)  
## S4 method for signature 'H2OModel'
h2o.confusionMatrix(object, newdata, valid = FALSE, ...)
## S4 method for signature 'H2OModelMetrics'
h2o.confusionMatrix(object, thresholds = NULL, metrics = NULL)
```

**Arguments**

- **object** Either an `H2OModel` object or an `H2OModelMetrics` object.
- **...** Extra arguments for extracting train or valid confusion matrices.
- **newdata** An H2OFrame object that can be scored on. Requires a valid response column.
- **valid** Retrieve the validation metric.
- **thresholds** (Optional) A value or a list of valid values between 0.0 and 1.0. This value is only used in the case of `H2OBinomialMetrics` objects.
- **metrics** (Optional) A metric or a list of valid metrics ("min_per_class_accuracy", "absolute_mcc", "tnr", "fmr", "fpr", "tpr", "precision", "accuracy", "f0point5", "f2", "f1"). This value is only used in the case of `H2OBinomialMetrics` objects.

**Details**

The `H2OModelMetrics` version of this function will only take `H2OBinomialMetrics` or `H2OMultinomialMetrics` objects. If no threshold is specified, all possible thresholds are selected.

**Value**

Calling this function on `H2OModel` objects returns a confusion matrix corresponding to the `predict` function. If used on an `H2OBinomialMetrics` object, returns a list of matrices corresponding to the number of thresholds specified.

**See Also**

- `predict` for generating prediction frames, `h2o.performance` for creating `H2OModelMetrics`. 
Examples

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(prostate_path)
prostate[, 2] <- as.factor(prostate[, 2])
model <- h2o.gbm(x = 3:9, y = 2, training_frame = prostate, distribution = "bernoulli")
h2o.confusionMatrix(model, prostate)
# Generating a ModelMetrics object
perf <- h2o.performance(model, prostate)
h2o.confusionMatrix(perf)

## End(Not run)
```

### h2o.connect

Connect to a running H2O instance.

**Description**

Connect to a running H2O instance.

**Usage**

```r
h2o.connect(
  ip = "localhost",
  port = 54321,
  strict_version_check = TRUE,
  proxy = NA_character_,
  https = FALSE,
  cacert = NA_character_,
  insecure = FALSE,
  username = NA_character_,
  password = NA_character_,
  use_spnego = FALSE,
  cookies = NA_character_,
  context_path = NA_character_,
  config = NULL
)
```

**Arguments**

- `ip` Object of class character representing the IP address of the server where H2O is running.
- `port` Object of class numeric representing the port number of the H2O server.
- `strict_version_check` (Optional) Setting this to FALSE is unsupported and should only be done when advised by technical support.
h2o.cor

Correlation of columns.

Description

Compute the correlation matrix of one or two H2OFrames.

Usage

```r
h2o.cor(x, y = NULL, na.rm = FALSE, use, method = "Pearson")
cor(x, ...)```
Arguments

\textbf{x} \hspace{1cm} \text{An H2OFrame object.}

\textbf{y} \hspace{1cm} \text{\texttt{NULL} (default) or an H2OFrame. The default is equivalent to \texttt{y = x}.}

\textbf{na.rm} \hspace{1cm} \text{\texttt{logical}. Should missing values be removed?}

\textbf{use} \hspace{1cm} \text{An optional character string indicating how to handle missing values. This must be one of the following: "everything" - outputs NaNs whenever one of its contributing observations is missing "all.obs" - presence of missing observations will throw an error "complete.obs" - discards missing values along with all observations in their rows so that only complete observations are used}

\textbf{method} \hspace{1cm} \text{\texttt{str} Method of correlation computation. Allowed values are: "Pearson" - Pearson's correlation coefficient "Spearman" - Spearman's correlation coefficient (Spearman's Rho) Defaults to "Pearson"}

\textbf{...} \hspace{1cm} \text{Further arguments to be passed down from other methods.}

Examples

```r
## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
cor(prostate$AGE)
## End(Not run)
```

---

\textbf{h2o.cos} \hspace{1cm} \textit{Compute the cosine of x}

Description

Compute the cosine of x

Usage

\texttt{h2o.cos(x)}

Arguments

\textbf{x} \hspace{1cm} \text{An H2OFrame object.}

See Also

\texttt{Trig} for the base R implementation, \texttt{cos()}. 
h2o.cosh

Examples

## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)
h2o.cos(frame["C1"])

## End(Not run)

h2o.cosh  Compute the hyperbolic cosine of x

Description

Compute the hyperbolic cosine of x

Usage

h2o.cosh(x)

Arguments

x  An H2OFrame object.

See Also

Hyperbolic for the base R implementation, cosh().

Examples

## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)
h2o.cosh(frame["C1"])

## End(Not run)
h2o.coxph

Trains a Cox Proportional Hazards Model (CoxPH) on an H2O dataset

Description

Trains a Cox Proportional Hazards Model (CoxPH) on an H2O dataset

Usage

h2o.coxph(
  x,
  event_column,
  training_frame,
  model_id = NULL,
  start_column = NULL,
  stop_column = NULL,
  weights_column = NULL,
  offset_column = NULL,
  stratify_by = NULL,
  ties = c("efron", "breslow"),
  init = 0,
  lre_min = 9,
  max_iterations = 20,
  interactions = NULL,
  interaction_pairs = NULL,
  interactions_only = NULL,
  use_all_factor_levels = FALSE,
  export_checkpoints_dir = NULL,
  single_node_mode = FALSE
)

Arguments

x (Optional) A vector containing the names or indices of the predictor variables to use in building the model. If x is missing, then all columns except event_column, start_column and stop_column are used.

event_column The name of binary data column in the training frame indicating the occurrence of an event.

training_frame Id of the training data frame.

model_id Destination id for this model; auto-generated if not specified.

start_column Start Time Column.

stop_column Stop Time Column.

weights_column Column with observation weights. Giving some observation a weight of zero is equivalent to excluding it from the dataset; giving an observation a relative
weight of 2 is equivalent to repeating that row twice. Negative weights are not allowed. Note: Weights are per-row observation weights and do not increase the size of the data frame. This is typically the number of times a row is repeated, but non-integer values are supported as well. During training, rows with higher weights matter more, due to the larger loss function pre-factor. If you set weight = 0 for a row, the returned prediction frame at that row is zero and this is incorrect. To get an accurate prediction, remove all rows with weight == 0.

offset_column Offset column. This will be added to the combination of columns before applying the link function.

stratify_by List of columns to use for stratification.

ties Method for Handling Ties. Must be one of: "efron", "breslow". Defaults to efron.

init Coefficient starting value. Defaults to 0.

lre_min Minimum log-relative error. Defaults to 9.

max_iterations Maximum number of iterations. Defaults to 20.

interactions A list of predictor column indices to interact. All pairwise combinations will be computed for the list.

interaction_pairs A list of pairwise (first order) column interactions.

interactions_only A list of columns that should only be used to create interactions but should not itself participate in model training.

use_all_factor_levels Logical. (Internal. For development only!) Indicates whether to use all factor levels. Defaults to FALSE.

export_checkpoints_dir Automatically export generated models to this directory.

single_node_mode Logical. Run on a single node to reduce the effect of network overhead (for smaller datasets) Defaults to FALSE.

Examples

```r
## Not run:
library(h2o)
h2o.init()

# Import the heart dataset
f <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/coxph_test/heart.csv"
heart <- h2o.importFile(f)

# Set the predictor and response
predictor <- "age"
response <- "event"

# Train a Cox Proportional Hazards model
heart_coxph <- h2o.coxph(x = predictor, training_frame = heart,
```
## Data H2OFrame Creation in H2O

### Description

Creates a data frame in H2O with real-valued, categorical, integer, and binary columns specified by the user.

### Usage

```r
h2o.createFrame(
  rows = 10000,
  cols = 10,
  randomize = TRUE,
  value = 0,
  real_range = 100,
  categorical_fraction = 0.2,
  factors = 100,
  integer_fraction = 0.2,
  integer_range = 100,
  binary_fraction = 0.1,
  binary_ones_fraction = 0.02,
  time_fraction = 0,
  string_fraction = 0,
  missing_fraction = 0.01,
  response_factors = 2,
  has_response = FALSE,
  seed,
  seed_for_column_types
)
```

### Arguments

- **rows**: The number of rows of data to generate.
- **cols**: The number of columns of data to generate. Excludes the response column if has_response = TRUE.
- **randomize**: A logical value indicating whether data values should be randomly generated. This must be TRUE if either categorical_fraction or integer_fraction is non-zero.
- **value**: If randomize = FALSE, then all real-valued entries will be set to this value.
- **real_range**: The range of randomly generated real values.
categorical_fraction
The fraction of total columns that are categorical.
factors
The number of (unique) factor levels in each categorical column.
integer_fraction
The fraction of total columns that are integer-valued.
integer_range
The range of randomly generated integer values.
binary_fraction
The fraction of total columns that are binary-valued.
binary_ones_fraction
The fraction of values in a binary column that are set to 1.
time_fraction
The fraction of randomly created date/time columns.
string_fraction
The fraction of randomly created string columns.
missing_fraction
The fraction of total entries in the data frame that are set to NA.
response_factors
If has_response = TRUE, then this is the number of factor levels in the response column.
has_response
A logical value indicating whether an additional response column should be prepended to the final H2O data frame. If set to TRUE, the total number of columns will be cols+1.
seed
A seed used to generate random values when randomize = TRUE.
seed_for_column_types
A seed used to generate random column types when randomize = TRUE.

Value

Returns an H2OFrame object.

Examples

```r
## Not run:
library(h2o)
h2o.init()
hf <- h2o.createFrame(rows = 1000, cols = 100, categorical_fraction = 0.1,
                       factors = 5, integer_fraction = 0.5, integer_range = 1,
                       has_response = TRUE)
head(hf)
summary(hf)

hf <- h2o.createFrame(rows = 100, cols = 10, randomize = FALSE, value = 5,
                       categorical_fraction = 0, integer_fraction = 0)
summary(hf)
## End(Not run)
```
h2o.cross_validation_fold_assignment

Retrieve the cross-validation fold assignment

Description

Retrieve the cross-validation fold assignment

Usage

h2o.cross_validation_fold_assignment(object)

Arguments

object An H2OModel object.

Value

Returns a H2OFrame

Examples

```r
## Not run:
library(h2o)
h2o.init()

cars <- h2o.importFile(f)
cars["economy_20mpg"] <- as.factor(cars["economy_20mpg"])
predictors <- c("displacement", "power", "weight", "acceleration", "year")
response <- "economy_20mpg"
cars_split <- h2o.splitFrame(data = cars, ratios = 0.8, seed = 1234)
train <- cars_split[[1]]
valid <- cars_split[[2]]
cars_gbm <- h2o.gbm(x = predictors, y = response, training_frame = train,
                   nfolds = 5, keep_cross_validation_fold_assignment = TRUE, seed = 1234)
h2o.cross_validation_fold_assignment(cars_gbm)

## End(Not run)
```
Retrieve the cross-validation holdout predictions

**Usage**

```r
h2o.cross_validation_holdout_predictions(object)
```

**Arguments**

- `object` An `H2OModel` object.

**Value**

Returns a H2OFrame

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
cars <- h2o.importFile(f)
cars["economy_20mpg"] <- as.factor(cars["economy_20mpg"])
predictors <- c("displacement","power","weight","acceleration","year")
response <- "economy_20mpg"
cars_split <- h2o.splitFrame(data = cars,ratios = 0.8, seed = 1234)
train <- cars_split[[1]]
valid <- cars_split[[2]]
cars_gbm <- h2o.gbm(x = predictors, y = response, training_frame = train,
nfolds = 5, keep_cross_validation_predictions = TRUE, seed = 1234)
h2o.cross_validation_holdout_predictions(cars_gbm)

## End(Not run)
```
h2o.cross_validation_models

Retrieves the cross-validation models

Description
Retrieve the cross-validation models

Usage
h2o.cross_validation_models(object)

Arguments
object An H2OModel object.

Value
Returns a list of H2OModel objects

Examples
## Not run:
library(h2o)
h2o.init()

cars <- h2o.importFile(f)
cars["economy_20mpg"] <- as.factor(cars["economy_20mpg"])
predictors <- c("displacement", "power", "weight", "acceleration", "year")
response <- "economy_20mpg"
cars_split <- h2o.splitFrame(data = cars, ratios = 0.8, seed = 1234)
train <- cars_split[[1]]
valid <- cars_split[[2]]
cars_gbm <- h2o.gbm(x = predictors, y = response, training_frame = train, 
nfolds = 5, keep_cross_validation_models = TRUE, seed = 1234)
h2o.cross_validation_models(cars_gbm)

## End(Not run)
h2o.cross_validation_predictions

Retrieve the cross-validation predictions

Description

Retrieve the cross-validation predictions

Usage

h2o.cross_validation_predictions(object)

Arguments

object An H2OModel object.

Value

Returns a list of H2OFrame objects

Examples

## Not run:
library(h2o)
h2o.init()

cars <- h2o.importFile(f)
cars["economy_20mpg"] <- as.factor(cars["economy_20mpg"])
predictors <- c("displacement", "power", "weight", "acceleration", "year")
response <- "economy_20mpg"
cars_split <- h2o.splitFrame(data = cars, ratios = 0.8, seed = 1234)
train <- cars_split[[1]]
valid <- cars_split[[2]]
cars_gbm <- h2o.gbm(x = predictors, y = response, training_frame = train,
                      nfolds = 5, keep_cross_validation_predictions = TRUE, seed = 1234)
h2o.cross_validation_predictions(cars_gbm)

## End(Not run)
h2o.cummax

*Return the cumulative max over a column or across a row*

### Description
Return the cumulative max over a column or across a row

### Usage

```r
h2o.cummax(x, axis = 0)
```

### Arguments
- `x`: An H2OFrame object.
- `axis`: An int that indicates whether to do down a column (0) or across a row (1).

### See Also
- `cumsum` for the base R implementation, `cummax()`.

### Examples

```r
## Not run:
library(h2o)
h2o.init()
frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)
h2o.cummax(frame, 1)
## End(Not run)
```

h2o.cummin

*Return the cumulative min over a column or across a row*

### Description
Return the cumulative min over a column or across a row

### Usage

```r
h2o.cummin(x, axis = 0)
```
Arguments

- `x`: An H2OFrame object.
- `axis`: An int that indicates whether to do down a column (0) or across a row (1).

See Also

cumsum for the base R implementation, cummin().

Examples

```r
## Not run:
library(h2o)
h2o.init()
frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)
h2o.cummin(frame, 1)
## End(Not run)
```

h2o.cumprod

Return the cumulative product over a column or across a row

Description

Return the cumulative product over a column or across a row

Usage

h2o.cumprod(x, axis = 0)

Arguments

- `x`: An H2OFrame object.
- `axis`: An int that indicates whether to do down a column (0) or across a row (1).

See Also

cumsum for the base R implementation, cumprod().
Examples

```r
## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)
h2o.cumprod(frame, 1)

## End(Not run)
```

h2o.cumsum

Return the cumulative sum over a column or across a row

Description

Return the cumulative sum over a column or across a row

Usage

```r
h2o.cumsum(x, axis = 0)
```

Arguments

- `x`: An H2OFrame object.
- `axis`: An int that indicates whether to do down a column (0) or across a row (1).

See Also

cumsum for the base R implementation.

Examples

```r
## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)
h2o.cumsum(frame, 1)

## End(Not run)
```
Description

Divides the range of the H2O data into intervals and codes the values according to which interval they fall in. The leftmost interval corresponds to the level one, the next is level two, etc.

Usage

h2o.cut(
  x,
  breaks,
  labels = NULL,
  include.lowest = FALSE,
  right = TRUE,
  dig.lab = 3,
  ...
)

## S3 method for class 'H2OFrame'
cut(
  x,
  breaks,
  labels = NULL,
  include.lowest = FALSE,
  right = TRUE,
  dig.lab = 3,
  ...
)

Arguments

- **x**: An H2OFrame object with a single numeric column.
- **breaks**: A numeric vector of two or more unique cut points.
- **labels**: Labels for the levels of the resulting category. By default, labels are constructed sing ":[(a,b]" interval notation.
- **include.lowest**: Logical, indicating if an 'x[j]' equal to the lowest (or highest, for right = FALSE 'breaks' value should be included.
- **right**: Logical, indicating if the intervals should be closed on the right (opened on the left) or vice versa.
- **dig.lab**: Integer which is used when labels are not given, determines the number of digits used in formatting the break numbers.
- **...**: Further arguments passed to or from other methods.
Value

Returns an H2OFrame object containing the factored data with intervals as levels.

Examples

```r
## Not run:
library(h2o)
h2o.init()
iris_hf <- as.h2o(iris)
summary(iris_hf)

# Cut sepal length column into intervals determined by min/max/quantiles
sepal_len_cut <- cut(iris_hf$Sepal.Length, c(4.2, 4.8, 5.8, 6, 8))
head(sepal_len_cut)
summary(sepal_len_cut)
## End(Not run)
```

**h2o.day**

*Convert Milliseconds to Day of Month in H2O Datasets*

**Description**

Converts the entries of an H2OFrame object from milliseconds to days of the month (on a 1 to 31 scale).

**Usage**

```r
h2o.day(x)
day(x)
```

```
## S3 method for class 'H2OFrame'
day(x)
```

**Arguments**

- `x` An H2OFrame object.

**Value**

An H2OFrame object containing the entries of `x` converted to days of the month.

**See Also**

- `h2o.month`
**h2o.dayOfWeek**

**Convert Milliseconds to Day of Week in H2O Datasets**

---

**Description**

Converts the entries of an H2OFrame object from milliseconds to days of the week (on a 0 to 6 scale).

**Usage**

```r
eh2o.dayOfWeek(x)
dayOfWeek(x)
```

**Arguments**

- `x` An H2OFrame object.

**Value**

An H2OFrame object containing the entries of `x` converted to days of the week.

**See Also**

- `h2o.day`, `h2o.month`

---

**h2o.dct**

**Compute DCT of an H2OFrame**

---

**Description**

Compute the Discrete Cosine Transform of every row in the H2OFrame.

**Usage**

```r
eh2o.dct(data, destination_frame, dimensions, inverse = FALSE)
```
**Arguments**

- `data`: An H2OFrame object representing the dataset to transform
- `destination_frame`: A frame ID for the result
- `dimensions`: An array containing the 3 integer values for height, width, depth of each sample. The product of HxWxD must total up to less than the number of columns. For 1D, use c(L,1,1), for 2D, use C(N,M,1).
- `inverse`: Whether to perform the inverse transform

**Value**

Returns an H2OFrame object.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
df <- h2o.createFrame(rows = 1000, cols = 8 * 16 * 24,
categorical_fraction = 0, integer_fraction = 0, missing_fraction = 0)
df1 <- h2o.dct(data = df, dimensions = c(8 * 16 * 24, 1, 1))
df2 <- h2o.dct(data = df1, dimensions = c(8 * 16 * 24, 1, 1), inverse = TRUE)
max(abs(df1 - df2))

df1 <- h2o.dct(data = df, dimensions = c(8 * 16, 24, 1))
df2 <- h2o.dct(data = df1, dimensions = c(8 * 16, 24, 1), inverse = TRUE)
max(abs(df1 - df2))

df1 <- h2o.dct(data = df, dimensions = c(8, 16, 24))
df2 <- h2o.dct(data = df1, dimensions = c(8, 16, 24), inverse = TRUE)
max(abs(df1 - df2))
## End(Not run)
```

---

**h2o.ddply**

*Split H2O Dataset, Apply Function, and Return Results*

**Description**

For each subset of an H2O data set, apply a user-specified function, then combine the results. This is an experimental feature based on plyr::ddply.

**Usage**

```r
h2o.ddply(X, .variables, FUN, ..., .progress = "none")
```
Arguments

- **X**: An H2OFrame object to be processed.
- **.variables**: Variables to split X by, either the indices or names of a set of columns.
- **FUN**: Function to apply to each subset grouping.
- **...**: Additional arguments passed on to FUN.
- **.progress**: Name of the progress bar to use. TODO: (Currently unimplemented)

Value

Returns an H2OFrame object containing the results from the split/apply operation, arranged

Examples

```r
## Not run:
library(h2o)
h2o.init()

# Import iris dataset to H2O
iris_hf <- as.h2o(iris)

# Add function taking mean of Sepal.Length column
fun <- function(df) { sum(df[, 1], na.rm = TRUE) / nrow(df) }

# Apply function to groups by flower specie
# uses h2o's `ddply, since iris_hf is an H2OFrame object
res <- h2o.ddply(iris_hf, "Species", fun)
head(res)

## End(Not run)
```

h2o.decryptionSetup

Setup a Decryption Tool

Description

If your source file is encrypted - setup a Decryption Tool and then provide the reference (result of this function) to the import functions.

Usage

```r
h2o.decryptionSetup(
  keystore,
  keystore_type = "JCEKS",
  key_alias = NA_character_,
  password = NA_character_,
  decrypt_tool = "",
  decrypt_impl = "water.parser.GenericDecryptionTool",
  cipher_spec = NA_character_)
```
h2o.deepfeatures

Feature Generation via H2O Deep Learning

Description

Extract the non-linear feature from an H2O data set using an H2O deep learning model.

Usage

h2o.deepfeatures(object, data, layer)

Arguments

object An H2OModel object that represents the deep learning model to be used for feature extraction.
data An H2OFrame object.
layer Index (integer) of the hidden layer to extract

h2o.deepfeatures

Feature Generation via H2O Deep Learning

Arguments

keystore An H2OFrame object referencing a loaded Java Keystore (see example).
keystore_type (Optional) Specification of Keystore type, defaults to JCEKS.
key_alias Which key from the keystore to use for decryption.
password Password to the keystore and the key.
decrypt_tool (Optional) Name of the decryption tool.
decrypt_impl (Optional) Java class name implementing the Decryption Tool.
cipher_spec Specification of a cipher (eg.: AES/ECB/PKCS5Padding).

See Also

h2o.importFile, h2o.parseSetup

Examples

```r
## Not run:
library(h2o)
h2o.init()
ks_path <- system.file("extdata", "keystore.jks", package = "h2o")
keystore <- h2o.importFile(path = ks_path, parse = FALSE) # don’t parse, keep as a binary file
cipher <- "AES/ECB/PKCS5Padding"
pwd <- "Password123"
alias <- "secretKeyAlias"
dt <- h2o.decryptionSetup(keystore, key_alias = alias, password = pwd, cipher_spec = cipher)
data_path <- system.file("extdata", "prostate.csv.aes", package = "h2o")
data <- h2o.importFile(data_path, decrypt_tool = dt)
summary(data)
## End(Not run)
```
Value

Returns an H2OFrame object with as many features as the number of units in the hidden layer of the specified index.

See Also

h2o.deeplearning for making H2O Deep Learning models.

Examples

```r
## Not run:
library(h2o)
h2o.init()
prostate_path = system.file("extdata", "prostate.csv", package = "h2o")
prostate = h2o.importFile(path = prostate_path)
prostate_dl = h2o.deeplearning(x = 3:9, y = 2, training_frame = prostate,
                              hidden = c(100, 200), epochs = 5)
prostate_deepfeatures_layer1 = h2o.deepfeatures(prostate_dl, prostate, layer = 1)
prostate_deepfeatures_layer2 = h2o.deepfeatures(prostate_dl, prostate, layer = 2)
head(prostate_deepfeatures_layer1)
head(prostate_deepfeatures_layer2)

## End(Not run)
```

---

h2o.deeplearning  

**Build a Deep Neural Network model using CPUs**

Description

Builds a feed-forward multilayer artificial neural network on an H2OFrame.

Usage

```r
h2o.deeplearning(
  x,
  y,
  training_frame,
  model_id = NULL,
  validation_frame = NULL,
  nfolds = 0,
  keep_cross_validation_models = TRUE,
  keep_cross_validation_predictions = FALSE,
  keep_cross_validation_fold_assignment = FALSE,
  fold_assignment = c("AUTO", "Random", "Modulo", "Stratified"),
  fold_column = NULL,
  ignore_const_cols = TRUE,
  score_each_iteration = FALSE,
)```
weights_column = NULL,
offset_column = NULL,
balance_classes = FALSE,
class_sampling_factors = NULL,
max_after_balance_size = 5,
checkpoint = NULL,
pretrained_autoencoder = NULL,
overwrite_with_best_model = TRUE,
use_all_factor_levels = TRUE,
standardize = TRUE,
activation = c("Tanh", "TanhWithDropout", "Rectifier", "RectifierWithDropout",
"Maxout", "MaxoutWithDropout"),
hidden = c(200, 200),
epochs = 10,
train_samples_per_iteration = -2,
target_ratio_comm_to_comp = 0.05,
seed = -1,
adaptive_rate = TRUE,
rho = 0.99,
epsilon = 1e-08,
rate = 0.005,
rate_annealing = 1e-06,
rate_decay = 1,
momentum_start = 0,
momentum_ramp = 1e+06,
momentum_stable = 0,
nesterov_accelerated_gradient = TRUE,
input_dropout_ratio = 0,
hidden_dropout_ratios = NULL,
l1 = 0,
l2 = 0,
max_w2 = 3.4028235e+38,
initial_weight_distribution = c("UniformAdaptive", "Uniform", "Normal"),
initial_weight_scale = 1,
initial_weights = NULL,
initial_biases = NULL,
loss = c("Automatic", "CrossEntropy", "Quadratic", "Huber", "Absolute", "Quantile"),
distribution = c("AUTO", "bernoulli", "multinomial", "gaussian", "poisson", "gamma",
"tweedie", "laplace", "quantile", "huber"),
quantile_alpha = 0.5,
tweedie_power = 1.5,
huber_alpha = 0.9,
score_interval = 5,
score_training_samples = 10000,
score_validation_samples = 0,
score_duty_cycle = 0.1,
classification_stop = 0,
regression_stop = 1e-06,
h2o.deeplearning

stopping_rounds = 5,
stopping_metric = c("AUTO", "deviance", "logloss", "MSE", "RMSE", "MAE", "RMSLE", "AUC", "AUCPR", "lift_top_group", "misclassification", "mean_per_class_error", "custom", "custom_increasing"),
stopping_tolerance = 0,
max_runtime_secs = 0,
score_validation_sampling = c("Uniform", "Stratified"),
diagnostics = TRUE,
fast_mode = TRUE,
force_load_balance = TRUE,
variable_importances = TRUE,
replicate_training_data = TRUE,
single_node_mode = FALSE,
shuffle_training_data = FALSE,
missing_values_handling = c("MeanImputation", "Skip"),
quiet_mode = FALSE,
autoencoder = FALSE,
sparse = FALSE,
col_major = FALSE,
average_activation = 0,
sparse_beta = 0,
max_categorical_features = 2147483647,
reproducible = FALSE,
export_weights_and_biases = FALSE,
mini_batch_size = 1,
elastic_averaging = FALSE,
elastic_averaging_moving_rate = 0.9,
elastic_averaging_regularization = 0.001,
export_checkpoints_dir = NULL,
auc_type = c("AUTO", "NONE", "MACRO_OVR", "WEIGHTED_OVR", "MACRO_OVO", "WEIGHTED_OVO"),
verbose = FALSE
)

Arguments

x (Optional) A vector containing the names or indices of the predictor variables to use in building the model. If x is missing, then all columns except y are used.

y The name or column index of the response variable in the data. The response must be either a numeric or a categorical/factor variable. If the response is numeric, then a regression model will be trained, otherwise it will train a classification model.

training_frame Id of the training data frame.

model_id Destination id for this model; auto-generated if not specified.

validation_frame Id of the validation data frame.
nfolds
  Number of folds for K-fold cross-validation (0 to disable or >= 2). Defaults to 0.

keep_cross_validation_models
  Logical. Whether to keep the cross-validation models. Defaults to TRUE.

keep_cross_validation_predictions
  Logical. Whether to keep the predictions of the cross-validation models. Defaults to FALSE.

keep_cross_validation_fold_assignment
  Logical. Whether to keep the cross-validation fold assignment. Defaults to FALSE.

fold_assignment
  Cross-validation fold assignment scheme, if fold_column is not specified. The 'Stratified' option will stratify the folds based on the response variable, for classification problems. Must be one of: "AUTO", "Random", "Modulo", "Stratified". Defaults to AUTO.

fold_column
  Column with cross-validation fold index assignment per observation.

ignore_const_cols
  Logical. Ignore constant columns. Defaults to TRUE.

score_each_iteration
  Logical. Whether to score during each iteration of model training. Defaults to FALSE.

weights_column
  Column with observation weights. Giving some observation a weight of zero is equivalent to excluding it from the dataset; giving an observation a relative weight of 2 is equivalent to repeating that row twice. Negative weights are not allowed. Note: Weights are per-row observation weights and do not increase the size of the data frame. This is typically the number of times a row is repeated, but non-integer values are supported as well. During training, rows with higher weights matter more, due to the larger loss function pre-factor. If you set weight = 0 for a row, the returned prediction frame at that row is zero and this is incorrect. To get an accurate prediction, remove all rows with weight == 0.

offset_column
  Offset column. This will be added to the combination of columns before applying the link function.

balance_classes
  Logical. Balance training data class counts via over/under-sampling (for imbalanced data). Defaults to FALSE.

class_sampling_factors
  Desired over/under-sampling ratios per class (in lexicographic order). If not specified, sampling factors will be automatically computed to obtain class balance during training. Requires balance_classes.

max_after_balance_size
  Maximum relative size of the training data after balancing class counts (can be less than 1.0). Requires balance_classes. Defaults to 5.0.

checkpoint
  Model checkpoint to resume training with.

pretrained_autoencoder
  Pretrained autoencoder model to initialize this model with.
overwrite_with_best_model
Logical. If enabled, override the final model with the best model found during training. Defaults to TRUE.

use_all_factor_levels
Logical. Use all factor levels of categorical variables. Otherwise, the first factor level is omitted (without loss of accuracy). Useful for variable importances and auto-enabled for autoencoder. Defaults to TRUE.

standardize
Logical. If enabled, automatically standardize the data. If disabled, the user must provide properly scaled input data. Defaults to TRUE.

activation

hidden
Hidden layer sizes (e.g. [100, 100]). Defaults to c(200, 200).

epochs
How many times the dataset should be iterated (streamed), can be fractional. Defaults to 10.

train_samples_per_iteration
Number of training samples (globally) per MapReduce iteration. Special values are 0: one epoch, -1: all available data (e.g., replicated training data), -2: automatic. Defaults to -2.

target_ratio_comm_to_comp
Target ratio of communication overhead to computation. Only for multi-node operation and train_samples_per_iteration = -2 (auto-tuning). Defaults to 0.05.

seed
Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Note: only reproducible when running single threaded. Defaults to -1 (time-based random number).

adaptive_rate
Logical. Adaptive learning rate. Defaults to TRUE.

rho
Adaptive learning rate time decay factor (similarity to prior updates). Defaults to 0.99.

epsilon
Adaptive learning rate smoothing factor (to avoid divisions by zero and allow progress). Defaults to 1e-08.

rate
Learning rate (higher => less stable, lower => slower convergence). Defaults to 0.005.

rate_annealing
Learning rate annealing: rate / (1 + rate_annealing * samples). Defaults to 1e-06.

rate_decay
Learning rate decay factor between layers (N-th layer: rate * rate_decay ^ (n - 1)). Defaults to 1.

momentum_start
Initial momentum at the beginning of training (try 0.5). Defaults to 0.

momentum_ramp
Number of training samples for which momentum increases. Defaults to 1000000.

momentum_stable
Final momentum after the ramp is over (try 0.99). Defaults to 0.

nesterov_accelerated_gradient
Logical. Use Nesterov accelerated gradient (recommended). Defaults to TRUE.

input_dropout_ratio
Input layer dropout ratio (can improve generalization, try 0.1 or 0.2). Defaults to 0.
hidden_dropout_ratios
  Hidden layer dropout ratios (can improve generalization), specify one value per
  hidden layer, defaults to 0.5.

l1
  L1 regularization (can add stability and improve generalization, causes many
  weights to become 0). Defaults to 0.

l2
  L2 regularization (can add stability and improve generalization, causes many
  weights to be small. Defaults to 0.

max_w2
  Constraint for squared sum of incoming weights per unit (e.g. for Rectifier).
  Defaults to 3.4028235e+38.

initial_weight_distribution
  Initial weight distribution. Must be one of: "UniformAdaptive", "Uniform",
  "Normal". Defaults to UniformAdaptive.

initial_weight_scale
  Uniform: -value...value, Normal: stddev. Defaults to 1.

initial_weights
  A list of H2OFrame ids to initialize the weight matrices of this model with.

initial_biases
  A list of H2OFrame ids to initialize the bias vectors of this model with.

loss
  Loss function. Must be one of: "Automatic", "CrossEntropy", "Quadratic", "Hub-
  ber", "Absolute", "Quantile". Defaults to Automatic.

distribution
  Distribution function Must be one of: "AUTO", "bernoulli", "multinomial",
  "gaussian", "poisson", "gamma", "tweedie", "laplace", "quantile", "huber". De-
 faults to AUTO.

quantile_alpha
  Desired quantile for Quantile regression, must be between 0 and 1. Defaults to
  0.5.

tweedie_power
  Tweedie power for Tweedie regression, must be between 1 and 2. Defaults to
  1.5.

huber_alpha
  Desired quantile for Huber/M-regression (threshold between quadratic and lin-
  ear loss, must be between 0 and 1). Defaults to 0.9.

score_interval
  Shortest time interval (in seconds) between model scoring. Defaults to 5.

score_training_samples
  Number of training set samples for scoring (0 for all). Defaults to 10000.

score_validation_samples
  Number of validation set samples for scoring (0 for all). Defaults to 0.

score_duty_cycle
  Maximum duty cycle fraction for scoring (lower: more training, higher: more
  scoring). Defaults to 0.1.

classification_stop
  Stopping criterion for classification error fraction on training data (-1 to disa-
  ble). Defaults to 0.

regression_stop
  Stopping criterion for regression error (MSE) on training data (-1 to disable).
  Defaults to 1e-06.

stopping_rounds
  Early stopping based on convergence of stopping_metric. Stop if simple moving
  average of length k of the stopping_metric does not improve for k:=stopping_rounds
  scoring events (0 to disable) Defaults to 5.
stopping_metric
Metric to use for early stopping (AUTO: logloss for classification, deviance for regression and anomaly_score for Isolation Forest). Note that custom and custom_increasing can only be used in GBM and DRF with the Python client. Must be one of: "AUTO", "deviance", "logloss", "MSE", "RMSE", "MAE", "RMSLE", "AUC", "AUCPR", "lift_top_group", "misclassification", "mean_per_class_error", "custom", "custom_increasing". Defaults to AUTO.

stopping_tolerance
Relative tolerance for metric-based stopping criterion (stop if relative improvement is not at least this much) Defaults to 0.

max_runtime_secs
Maximum allowed runtime in seconds for model training. Use 0 to disable. Defaults to 0.

score_validation_sampling
Method used to sample validation dataset for scoring. Must be one of: "Uniform", "Stratified". Defaults to Uniform.

diagnostics
Logical. Enable diagnostics for hidden layers. Defaults to TRUE.

fast_mode
Logical. Enable fast mode (minor approximation in back-propagation). Defaults to TRUE.

force_load_balance
Logical. Force extra load balancing to increase training speed for small datasets (to keep all cores busy). Defaults to TRUE.

variable_importances
Logical. Compute variable importances for input features (Gedeon method) - can be slow for large networks. Defaults to TRUE.

replicate_training_data
Logical. Replicate the entire training dataset onto every node for faster training on small datasets. Defaults to TRUE.

single_node_mode
Logical. Run on a single node for fine-tuning of model parameters. Defaults to FALSE.

shuffle_training_data
Logical. Enable shuffling of training data (recommended if training data is replicated and train_samples_per_iteration is close to #nodes x #rows, of if using balance_classes). Defaults to FALSE.

missing_values_handling
Handling of missing values. Either MeanImputation or Skip. Must be one of: "MeanImputation", "Skip". Defaults to MeanImputation.

quiet_mode
Logical. Enable quiet mode for less output to standard output. Defaults to FALSE.

autoencoder
Logical. Auto-Encoder. Defaults to FALSE.

sparse
Logical. Sparse data handling (more efficient for data with lots of 0 values). Defaults to FALSE.

col_major
Logical. DEPRECATED Use a column major weight matrix for input layer. Can speed up forward propagation, but might slow down backpropagation. Defaults to FALSE.
average_activation
Average activation for sparse auto-encoder. #Experimental Defaults to 0.

sparsity_beta
Sparsity regularization. #Experimental Defaults to 0.

max_categorical_features
Max. number of categorical features, enforced via hashing. #Experimental Defaults to 2147483647.

reproducible
Logical. Force reproducibility on small data (will be slow - only uses 1 thread). Defaults to FALSE.

export_weights_and_biases
Logical. Whether to export Neural Network weights and biases to H2O Frames. Defaults to FALSE.

mini_batch_size
Mini-batch size (smaller leads to better fit, larger can speed up and generalize better). Defaults to 1.

categorical_encoding
Encoding scheme for categorical features Must be one of: "AUTO", "Enum", "OneHotInternal", "OneHotExplicit", "Binary", "Eigen", "LabelEncoder", "Sort-ByResponse", "EnumLimited". Defaults to AUTO.

elastic_averaging
Logical. Elastic averaging between compute nodes can improve distributed model convergence. #Experimental Defaults to FALSE.

elastic_averaging_moving_rate
Elastic averaging moving rate (only if elastic averaging is enabled). Defaults to 0.9.

elastic_averaging_regularization
Elastic averaging regularization strength (only if elastic averaging is enabled). Defaults to 0.001.

export_checkpoints_dir
Automatically export generated models to this directory.

auc_type
Set default multinomial AUC type. Must be one of: "AUTO", "NONE", "MACRO_OVR", "WEIGHTED_OVR", "MACRO_OVO", "WEIGHTED_OVO". Defaults to AUTO.

verbose
Logical. Print scoring history to the console (Metrics per epoch). Defaults to FALSE.

See Also
predict.H2OModel for prediction

Examples

```r
# Not run:
library(h2o)
h2o.init()
iris_hf <- as.h2o(iris)
iris_dl <- h2o.deeplearning(x = 1:4, y = 5, training_frame = iris_hf, seed=123456)

# now make a prediction
predictions <- h2o.predict(iris_dl, iris_hf)
```
h2o.describe

### H2O Description of A Dataset

**Description**

Reports the "Flow" style summary rollups on an instance of H2OFrame. Includes information about column types, mins/maxs/missing/zero counts/stds/number of levels

**Usage**

h2o.describe(frame)

**Arguments**

- frame: An H2OFrame object.

**Value**

A table with the Frame stats.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.importFile(path = prostate_path)
h2o.describe(prostate)

## End(Not run)
```

h2o.difflag1

### Conduct a lag 1 transform on a numeric H2OFrame column

**Description**

Conduct a lag 1 transform on a numeric H2OFrame column

**Usage**

```r
h2o.difflag1(object)
```

**Arguments**

```r
- object: H2OFrame object
```
h2o.dim

Returns the number of rows and columns for an H2OFrame object.

Description

Returns the number of rows and columns for an H2OFrame object.

Usage

h2o.dim(x)

Arguments

x

An H2OFrame object.

See Also

dim for the base R implementation.

Examples

## Not run:
library(h2o)
h2o.init()

cars <- h2o.importFile(f)
predictors <- c("displacement", "power", "weight", "acceleration", "year")
response <- "cylinders"
cars_split <- h2o.splitFrame(data = cars, ratios = 0.8, seed = 1234)
train <- cars_split[[1]]
valid <- cars_split[[2]]
cars_gbm <- h2o.gbm(x = predictors, y = response, training_frame = train,
validation_frame = valid, nfolds = 5, seed = 1234)
h2o.difflag1(cars["cylinders"])

## End(Not run)
h2o.dimnames

Column names of an H2OFrame

Description

Column names of an H2OFrame

Usage

h2o.dimnames(x)

Arguments

x
An H2OFrame object.

See Also

dimnames for the base R implementation.

Examples

## Not run:
library(h2o)
h2o.init()

cars <- h2o.importFile(f)
h2o.dimnames(cars)

## End(Not run)

h2o.distance

Compute a pairwise distance measure between all rows of two numeric H2OFrames.

Description

Compute a pairwise distance measure between all rows of two numeric H2OFrames.

Usage

h2o.distance(x, y, measure)
Arguments

- **x**: An H2OFrame object (large, references).
- **y**: An H2OFrame object (small, queries).
- **measure**: An optional string indicating what distance measure to use. Must be one of: "l1" - Absolute distance (L1-norm, >=0) "l2" - Euclidean distance (L2-norm, >=0) "cosine" - Cosine similarity (-1...1) "cosine_sq" - Squared Cosine similarity (0...1)

Examples

```r
## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
h2o.distance(prostate[11:30, ], prostate[1:10, ], "cosine")
## End(Not run)
```

---

**h2o.downloadAllLogs**

Download H2O Log Files to Disk

Description

h2o.downloadAllLogs downloads all H2O log files to local disk in .zip format. Generally used for debugging purposes.

Usage

```r
h2o.downloadAllLogs(dirname = ".", filename = NULL)
```

Arguments

- **dirname**: (Optional) A character string indicating the directory that the log file should be saved in.
- **filename**: (Optional) A character string indicating the name that the log file should be saved to. Note that the saved format is .zip, so the file name must include the .zip extension.

Examples

```r
## Not run:
h2o.downloadAllLogs(dirname='./your_directory_name/', filename = 'autoh2o_log.zip')
## End(Not run)
```
**Description**

Download an H2O data set to a CSV file on the local disk

**Usage**

\[
\text{h2o.downloadCSV}(\text{data}, \text{filename})
\]

**Arguments**

- **data**: an H2OFrame object to be downloaded.
- **filename**: A string indicating the name that the CSV file should be saved to.

**Warning**

Files located on the H2O server may be very large! Make sure you have enough hard drive space to accommodate the entire file.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
iris_hf <- as.h2o(iris)

file_path <- paste(getwd(), "my_iris_file.csv", sep = .Platform$file.sep)
h2o.downloadCSV(iris_hf, file_path)
file.info(file_path)
file.remove(file_path)

## End(Not run)
```

**Description**

Download the model in binary format. The owner of the file saved is the user by which python session was executed.
Usage

h2o.download_model(
    model,
    path = NULL,
    export_cross_validation_predictions = FALSE,
    filename = ""
)

Arguments

model An H2OModel
path The path where binary file should be downloaded. Downloaded to current directory by default.
export_cross_validation_predictions A boolean flag indicating whether the download model should be saved with CV Holdout Frame predictions. Default is not to export the predictions.
filename string indicating the file name.

Examples

## Not run:
library(h2o)
h <- h2o.init()
fr <- as.h2o(iris)
my_model <- h2o.gbm(x = 1:4, y = 5, training_frame = fr)
h2o.download_model(my_model) # save to the current working directory

## End(Not run)

h2o.download_mojo

Download the model in MOJO format.

Usage

h2o.download_mojo(
    model,
    path = getwd(),
    get_genmodel_jar = FALSE,
    genmodel_name = "",
    genmodel_path = "",
    filename = ""
)
Arguments

model: An H2OModel
path: The path where MOJO file should be saved. Saved to current directory by default.
get_genmodel_jar: If TRUE, then also download h2o-genmodel.jar and store it in either the same folder as the MOJO or in “genmodel_path” if specified.
genmodel_name: Custom name of genmodel jar.
genmodel_path: Path to store h2o-genmodel.jar. If left blank and “get_genmodel_jar” is TRUE, then the h2o-genmodel.jar is saved to “path”.
filename: string indicating the file name. (Type of file is always .zip)

Value

Name of the MOJO file written to the path.

Examples

```r
## Not run:
library(h2o)
h <- h2o.init()
fr <- as.h2o(iris)
my_model <- h2o.gbm(x = 1:4, y = 5, training_frame = fr)
h2o.download_mojo(my_model) # save to the current working directory
## End(Not run)
```

h2o.download_pojo

**Download the Scoring POJO (Plain Old Java Object) of an H2O Model**

Description

Download the Scoring POJO (Plain Old Java Object) of an H2O Model

Usage

```r
h2o.download_pojo(
  model, 
  path = NULL, 
  getjar = NULL, 
  get_jar = TRUE, 
  jar_name = ""
)
```
h2o.drop_duplicates

Drops duplicated rows.

**Description**

Drops duplicated rows across specified columns.

**Usage**

```r
h2o.drop_duplicates(frame, columns, keep = "first")
```

**Arguments**

- `frame`: An H2OFrame object to drop duplicates on.
- `columns`: Columns to compare during the duplicate detection process.
- `keep`: Which rows to keep. The "first" value (default) keeps the first row and deletes the rest. The "last" keeps the last row.
h2o.entropy

### Not run:
```r
library(h2o)
h2o.init()

data <- as.h2o(iris)
deduplicated_data <- h2o.drop_duplicates(data, c("Species", "Sepal.Length"), keep = "first")
```

### Shannon entropy

**Description**

Return the Shannon entropy of a string column. If the string is empty, the entropy is 0.

**Usage**

```r
h2o.entropy(x)
```

**Arguments**

- `x`: The column on which to calculate the entropy.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

buys <- as.h2o(c("no", "no", "yes", "yes", "yes", "no", "no", "yes", "yes", "yes", "no"))
buys_entropy <- h2o.entropy(buys)
```

h2o.exp

### Compute the exponential function of x

**Description**

Compute the exponential function of x

**Usage**

```r
h2o.exp(x)
```
Arguments

- x: An H2OFrame object.

See Also

- Log for the base R implementation, exp()

Examples

```r
## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)
h2o.exp(frame["C1"])
## End(Not run)
```

---

**h2o.explain**

*Generate Model Explanations*

**Description**

The H2O Explainability Interface is a convenient wrapper to a number of explainability methods and visualizations in H2O. The function can be applied to a single model or group of models and returns a list of explanations, which are individual units of explanation such as a partial dependence plot or a variable importance plot. Most of the explanations are visual (ggplot plots). These plots can also be created by individual utility functions as well.

**Usage**

```r
h2o.explain(
  object,  # H2O model
  newdata, # Optional new data
  columns = NULL, # Optional column indices
  top_n_features = 5, # Number of top features
  include_explanations = "ALL", # Include explanations
  exclude_explanations = NULL, # Exclude explanations
  plot_overrides = NULL # Plot overrides
)
```
Arguments

- **object**: A list of H2O models, an H2O AutoML instance, or an H2OFrame with a 'model_id' column (e.g. H2OAutoML leaderboard).
- **newdata**: An H2OFrame.
- **columns**: A vector of column names or column indices to create plots with. If specified parameter top_n_features will be ignored.
- **top_n_features**: An integer specifying the number of columns to use, ranked by variable importance (where applicable).
- **include_explanations**: If specified, return only the specified model explanations. (Mutually exclusive with exclude_explanations)
- **exclude_explanations**: Exclude specified model explanations.
- **plot_overrides**: Overrides for individual model explanations, e.g. list(shap_summary_plot = list(columns = 50)).

Value

List of outputs with class "H2OExplanation"

Examples

```r
## Not run:
library(h2o)
h2o.init()

# Import the wine dataset into H2O:
df <- h2o.importFile(f)

# Set the response
response <- "quality"

# Split the dataset into a train and test set:
splits <- h2o.splitFrame(df, ratios = 0.8, seed = 1)
train <- splits[[1]]
test <- splits[[2]]

# Build and train the model:
aml <- h2o.automl(y = response,
                 training_frame = train,
                 max_models = 10,
                 seed = 1)

# Create the explanation for whole H2OAutoML object
exa <- h2o.explain(aml, test)
print(exa)

# Create the explanation for the leader model
```
exm <- h2o.explain(aml@leader, test)
print(exm)

## End(Not run)

h2o.explain_row

Generate Model Explanations for a single row

Description

Explain the behavior of a model or group of models with respect to a single row of data. The function returns a list of explanations, which are individual units of explanation such as a partial dependence plot or a variable importance plot. Most of the explanations are visual (ggplot plots). These plots can also be created by individual utility functions as well.

Usage

h2o.explain_row(
  object,
  newdata,
  row_index,
  columns = NULL,
  top_n_features = 5,
  include_explanations = "ALL",
  exclude_explanations = NULL,
  plot_overrides = NULL
)

Arguments

object A list of H2O models, an H2O AutoML instance, or an H2OFrame with a 'model_id' column (e.g. H2OAutoML leaderboard).
newdata An H2OFrame.
row_index A row index of the instance to explain.
columns A vector of column names or column indices to create plots with. If specified parameter top_n_features will be ignored.
top_n_features An integer specifying the number of columns to use, ranked by variable importance (where applicable).
include_explanations If specified, return only the specified model explanations. (Mutually exclusive with exclude_explanations)
exclude_explanations Exclude specified model explanations.
plot_overrides Overrides for individual model explanations, e.g., list(shap_explain_row = list(columns = 5))
Value

List of outputs with class "H2OExplanation"

Examples

```r
## Not run:
library(h2o)
h2o.init()

# Import the wine dataset into H2O:
df <- h2o.importFile(f)

# Set the response
response <- "quality"

# Split the dataset into a train and test set:
splits <- h2o.splitFrame(df, ratios = 0.8, seed = 1)
train <- splits[[1]]
test <- splits[[2]]

# Build and train the model:
aml <- h2o.automl(y = response,
                   training_frame = train,
                   max_models = 10,
                   seed = 1)

# Create the explanation for whole H2OAutoML object
exa <- h2o.explain_row(aml, test, row_index = 1)
print(exa)

# Create the explanation for the leader model
exm <- h2o.explain_row(aml@leader, test, row_index = 1)
print(exm)

## End(Not run)
```

h2o.exportFile

Export an H2O Data Frame (H2OFrame) to a File or to a collection of Files.

Description

Exports an H2OFrame (which can be either VA or FV) to a file. This file may be on the H2O instance’s local filesystem, or to HDFS (preface the path with hdfs://) or to S3N (preface the path with s3n://).
Usage

h2o.exportFile(
  data,
  path,
  force = FALSE,
  sep = ",",
  compression = NULL,
  parts = 1,
  header = TRUE,
  quote_header = TRUE
)

Arguments

data An H2OFrame object.

path The path to write the file to. Must include the directory and also filename if exporting to a single file. May be prefaced with hdfs:// or s3n://. Each row of data appears as line of the file.

force logical, indicates how to deal with files that already exist.

sep The field separator character. Values on each line of the file will be separated by this character (default ",").

compression How to compress the exported dataset (default none; gzip, bzip2 and snappy available)

parts integer, number of part files to export to. Default is to write to a single file. Large data can be exported to multiple 'part' files, where each part file contains subset of the data. User can specify the maximum number of part files or use value -1 to indicate that H2O should itself determine the optimal number of files. Parameter path will be considered to be a path to a directory if export to multiple part files is desired. Part files conform to naming scheme 'part-m-?????'.

header logical, indicates whether to write the header line. Default is to include the header in the output file.

quote_header logical, indicates whether column names should be quoted. Default is to use quotes.

Details

In the case of existing files force = TRUE will overwrite the file. Otherwise, the operation will fail.

Examples

## Not run:
library(h2o)
h2o.init()
iris_hf <- as.h2o(iris)

# These aren't real paths
# h2o.exportFile(iris_hf, path = "/path/on/h2o/server/filesystem/iris.csv")
h2o.exportHDFS

Export a Model to HDFS

Description

Exports an H2OModel to HDFS.

Usage

h2o.exportHDFS(object, path, force = FALSE)

Arguments

- **object**: an H2OModel class object.
- **path**: The path to write the model to. Must include the directory and filename.
- **force**: logical, indicates how to deal with files that already exist.

Examples

```r
## Not run:
library(h2o)
h2o.init

train <- h2o.importFile(f)
h2o.exportHDFS(train, path = "", force = FALSE)

## End(Not run)
```

h2o.extendedIsolationForest

Trains an Extended Isolation Forest model

Description

Trains an Extended Isolation Forest model
Usage

h2o.extendedIsolationForest(
  training_frame,
  x,
  model_id = NULL,
  ignore_const_cols = TRUE,
  categorical_encoding = c("AUTO", "Enum", "OneHotInternal", "OneHotExplicit",
    "Binary", "Eigen", "LabelEncoder", "SortByResponse", "EnumLimited"),
  ntrees = 100,
  sample_size = 256,
  extension_level = 0,
  seed = -1
)

Arguments

training_frame  Id of the training data frame.

x  A vector containing the character names of the predictors in the model.

model_id  Destination id for this model; auto-generated if not specified.

ignore_const_cols  Logical. Ignore constant columns. Defaults to TRUE.

categorical_encoding  Encoding scheme for categorical features Must be one of: "AUTO", "Enum", "OneHotInternal", "OneHotExplicit", "Binary", "Eigen", "LabelEncoder", "SortByResponse", "EnumLimited". Defaults to AUTO.

ntrees  Number of Extended Isolation Forest trees. Defaults to 100.

sample_size  Number of randomly sampled observations used to train each Extended Isolation Forest tree. Defaults to 256.

extension_level  Maximum is N - 1 (N = numCols). Minimum is 0. Extended Isolation Forest with extension_level = 0 behaves like Isolation Forest. Defaults to 0.

seed  Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).

Examples

## Not run:
library(h2o)
h2o.init()

# Import the prostate dataset
p <- h2o.importFile(path="https://raw.github.com/h2oai/h2o/master/smalldata/logreg/prostate.csv")

# Set the predictors
predictors <- c(\"AGE\", \"RACE\", \"DPROS\", \"DCAPS\", \"PSA\", \"VOL\", \"GLEASON\")
# Build an Extended Isolation forest model
model <- h2o.extendedIsolationForest(x = predictors,
                                      training_frame = p,
                                      model_id = "eif.hex",
                                      ntrees = 100,
                                      sample_size = 256,
                                      extension_level = length(predictors) - 1)

# Calculate score
score <- h2o.predict(model, p)
anomaly_score <- score$anomaly_score

# Number in [0, 1] explicitly defined in Equation (1) from Extended Isolation Forest paper
# or in paragraph '2 Isolation and Isolation Trees' of Isolation Forest paper
anomaly_score <- score$anomaly_score

# Average path length of the point in Isolation Trees from root to the leaf
mean_length <- score$mean_length

## End(Not run)

---

**h2o.feature_interaction**

*Feature interactions and importance, leaf statistics and split value histograms in a tabular form. Available for XGBoost and GBM.*

**Description**

Metrics: Gain - Total gain of each feature or feature interaction. FScore - Amount of possible splits taken on a feature or feature interaction. wFScore - Amount of possible splits taken on a feature or feature interaction weighted by the probability of the splits to take place. Average wFScore - wFScore divided by FScore. Average Gain - Gain divided by FScore. Expected Gain - Total gain of each feature or feature interaction weighted by the probability to gather the gain. Average Tree Index Average Tree Depth

**Usage**

```r
h2o.feature_interaction(
  model,
  max_interaction_depth = 100,
  max_tree_depth = 100,
  max_deepening = -1
)
```

**Arguments**

- `model` A trained xgboost model.
- `max_interaction_depth` Upper bound for extracted feature interactions depth. Defaults to 100.

max_tree_depth  Upper bound for tree depth. Defaults to 100.
max_deepeening  Upper bound for interaction start deepening (zero deepening => interactions starting at root only). Defaults to -1.

Examples

```r
## Not run:
library(h2o)
h2o.init()
boston <- h2o.importFile(
  destination_frame="boston"
)
boston_xgb <- h2o.xgboost(training_frame = boston, y = "medv", seed = 1234)
feature_interactions <- h2o.feature_interaction(boston_xgb)

## End(Not run)
```

h2o.fillna

```
fillNA
```

Description

Fill NA's in a sequential manner up to a specified limit

Usage

```r
h2o.fillna(x, method = "forward", axis = 1, maxlen = 1L)
```

Arguments

- `x`: an H2OFrame
- `method`: A String: "forward" or "backward"
- `axis`: An Integer 1 for row-wise fill (default), 2 for column-wise fill
- `maxlen`: An Integer for maximum number of consecutive NA's to fill

Value

An H2OFrame after filling missing values

Examples

```r
## Not run:
library(h2o)
h2o.init()

frame_with_nas <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
```
h2o.filterNACols

Description

Filter NA Columns

Usage

h2o.filterNACols(data, frac = 0.2)

Arguments

data
  A dataset to filter on.
frac
  The threshold of NAs to allow per column (columns >= this threshold are filtered)

Value

Returns a numeric vector of indexes that pertain to non-NA columns

Examples

## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)
h2o.filterNACols(frame, frac = 0.5)
h2o.filterNACols(frame, frac = 0.6)

## End(Not run)
h2o.findSynonyms  
Find synonyms using a word2vec model.

Description
Find synonyms using a word2vec model.

Usage
h2o.findSynonyms(word2vec, word, count = 20)

Arguments
- word2vec: A word2vec model.
- word: A single word to find synonyms for.
- count: The top ‘count’ synonyms will be returned.

Examples
## Not run:
library(h2o)
h2o.init()

job_titles <- h2o.importFile(
  col.names = c("category", "jobtitle"), col.types = c("String", "String"), header = TRUE)
words <- h2o.tokenize(job_titles, " ")
vec <- h2o.word2vec(training_frame = words)
h2o.findSynonyms(vec, "teacher", count = 20)

## End(Not run)

h2o.find_row_by_threshold  
Find the threshold, give the max metric. No duplicate thresholds allowed.

Description
Find the threshold, give the max metric. No duplicate thresholds allowed.

Usage
h2o.find_row_by_threshold(object, threshold)
**h2o.find_threshold_by_max_metric**

*Find the threshold, give the max metric*

### Description

Find the threshold, give the max metric

### Usage

```r
h2o.find_threshold_by_max_metric(object, metric)
```

### Arguments

- `object` H2OBinomialMetrics
- `metric` "F1," for example
Examples

```r
## Not run:
library(h2o)
h2o.init()

cars <- h2o.importFile(f)
cars["economy_20mpg"] <- as.factor(cars["economy_20mpg"])
predictors <- c("displacement", "power", "weight", "acceleration", "year")
response <- "economy_20mpg"
cars_split <- h2o.splitFrame(data = cars, ratios = 0.8, seed = 1234)
train <- cars_split[[1]]
valid <- cars_split[[2]]
cars_gbm <- h2o.gbm(x = predictors, y = response,
  training_frame = train, validation_frame = valid,
  build_tree_one_node = TRUE, seed = 1234)
perf <- h2o.performance(cars_gbm, cars)
h2o.find_threshold_by_max_metric(perf, "fnr")

## End(Not run)
```

---

**h2o.floor**

*Take a single numeric argument and return a numeric vector with the largest integers*

Description

`floor` takes a single numeric argument `x` and returns a numeric vector containing the largest integers not greater than the corresponding elements of `x`.

Usage

```r
h2o.floor(x)
```

Arguments

- `x` An H2OFrame object.

See Also

`Round` for the base R implementation, `floor()`.

Examples

```r
## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
```
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)

h2o.floor(frame["C2"])

## End(Not run)

---

### h2o.flow

*Open H2O Flow*

**Description**

Open H2O Flow in your browser

**Usage**

h2o.flow()

---

### h2o.gainsLift

*Access H2O Gains/Lift Tables*

**Description**

Retrieve either a single or many Gains/Lift tables from H2O objects.

**Usage**

h2o.gainsLift(object, ...)

```r
## S4 method for signature 'H2OModel'

h2o.gainsLift(object, newdata, valid = FALSE, xval = FALSE, ...)
```

```r
## S4 method for signature 'H2OModelMetrics'

h2o.gainsLift(object)
```

**Arguments**

- **object**: Either an `H2OModel` object or an `H2OModelMetrics` object.
- **...**: further arguments to be passed to/from this method.
- **newdata**: An H2OFrame object that can be scored on. Requires a valid response column.
- **valid**: Retrieve the validation metric.
- **xval**: Retrieve the cross-validation metric.

**Details**

The `H2OModelMetrics` version of this function will only take `H2OBinomialMetrics` objects.
Value

Calling this function on H2OModel objects returns a Gains/Lift table corresponding to the predict function.

See Also

predict for generating prediction frames, h2o.performance for creating H2OModelMetrics.

Examples

## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(prostate_path)
prostate[, 2] <- as.factor(prostate[, 2])
model <- h2o.gbm(x = 3:9, y = 2, distribution = "bernoulli",
                 training_frame = prostate, validation_frame = prostate, nfolds = 3)
h2o.gainsLift(model) ## extract training metrics
h2o.gainsLift(model, valid = TRUE) ## extract validation metrics (here: the same)
h2o.gainsLift(model, xval = TRUE) ## extract cross-validation metrics
h2o.gainsLift(model, newdata = prostate) ## score on new data (here: the same)
# Generating a ModelMetrics object
perf <- h2o.performance(model, prostate)
h2o.gainsLift(perf) ## extract from existing metrics object

## End(Not run)

h2o.gam

Fit a General Additive Model

Description

Creates a generalized additive model, specified by a response variable, a set of predictors, and a description of the error distribution.

Usage

h2o.gam(
  x,
  y,
  training_frame,
  gam_columns,
  model_id = NULL,
  validation_frame = NULL,
  nfolds = 0,
  seed = -1,
  keep_cross_validation_models = TRUE,
)
keep_cross_validation_predictions = FALSE,
keep_cross_validation_fold_assignment = FALSE,
fold_assignment = c("AUTO", "Random", "Modulo", "Stratified"),
fold_column = NULL,
ignore_const_cols = TRUE,
score_each_iteration = FALSE,
offset_column = NULL,
weights_column = NULL,
family = c("AUTO", "gaussian", "binomial", "quasibinomial", "ordinal", "multinomial",
          "poisson", "gamma", "tweedie", "negativebinomial", "fractionalbinomial"),
tweedie_variance_power = 0,
tweedie_link_power = 0,
theta = 0,
solver = c("AUTO", "IRLSM", "L_BFGS", "COORDINATE_DESCENT_NAIVE",
          "COORDINATE_DESCENT", "GRADIENT_DESCENT_LH", "GRADIENT_DESCENT_SQERR"),
alPHA = NULL,
lambda = NULL,
lambda_search = FALSE,
early_stopping = TRUE,
nlambdas = -1,
standardize = FALSE,
missing_values_handling = c("MeanImputation", "Skip", "PlugValues"),
plug_values = NULL,
compute_p_values = FALSE,
remove_collinear_columns = FALSE,
intercept = TRUE,
non_negative = FALSE,
max_iterations = -1,
objective_epsilon = -1,
beta_epsilon = 1e-04,
gradient_epsilon = -1,
link = c("family_default", "identity", "logit", "log", "inverse", "tweedie",
         "ologit"),
startval = NULL,
prior = -1,
cold_start = FALSE,
lambda_min_ratio = -1,
beta_constraints = NULL,
max_active_predictors = -1,
interactions = NULL,
interaction_pairs = NULL,
obj_reg = -1,
export_checkpoints_dir = NULL,
stopping_rounds = 0,
stopping_metric = c("AUTO", "deviance", "logloss", "MSE", "RMSE", "MAE", "RMSLE",
                    "AUC", "AUCPR", "lift_top_group", "misclassification", "mean_per_class_error",
                    "custom", "custom_increasing"),
stopping_tolerance = 0.001,
Arguments

x (Optional) A vector containing the names or indices of the predictor variables to use in building the model. If x is missing, then all columns except y are used.

y The name or column index of the response variable in the data. The response must be either a numeric or a categorical/factor variable. If the response is numeric, then a regression model will be trained, otherwise it will train a classification model.

training_frame Id of the training data frame.

gam_columns Arrays of predictor column names for gam for smoothers using single or multiple predictors like 'c1','c2','c3','c4',....

model_id Destination id for this model; auto-generated if not specified.

validation_frame Id of the validation data frame.

nfolds Number of folds for K-fold cross-validation (0 to disable or >= 2). Defaults to 0.

seed Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).

keep_cross_validation_models Logical. Whether to keep the cross-validation models. Defaults to TRUE.

keep_cross_validation_predictions Logical. Whether to keep the predictions of the cross-validation models. Defaults to FALSE.

keep_cross_validation_fold_assignment Logical. Whether to keep the cross-validation fold assignment. Defaults to FALSE.

fold_assignment Cross-validation fold assignment scheme, if fold_column is not specified. The 'Stratified' option will stratify the folds based on the response variable, for classification problems. Must be one of: "AUTO", "Random", "Modulo", "Stratified". Defaults to AUTO.
fold_column  Column with cross-validation fold index assignment per observation.
ignore_const_cols Logical. Ignore constant columns. Defaults to TRUE.
score_each_iteration Logical. Whether to score during each iteration of model training. Defaults to FALSE.
offset_column Offset column. This will be added to the combination of columns before applying the link function.
weights_column Column with observation weights. Giving some observation a weight of zero is equivalent to excluding it from the dataset; giving an observation a relative weight of 2 is equivalent to repeating that row twice. Negative weights are not allowed. Note: Weights are per-row observation weights and do not increase the size of the data frame. This is typically the number of times a row is repeated, but non-integer values are supported as well. During training, rows with higher weights matter more, due to the larger loss function pre-factor. If you set weight = 0 for a row, the returned prediction frame at that row is zero and this is incorrect. To get an accurate prediction, remove all rows with weight == 0.
family Family. Use binomial for classification with logistic regression, others are for regression problems. Must be one of: "AUTO", "gaussian", "binomial", "quasibinomial", "ordinal", "multinomial", "poisson", "gamma", "tweedie", "negativebinomial", "fractionalbinomial". Defaults to AUTO.
tweedie_variance_power Tweedie variance power. Defaults to 0.
tweedie_link_power Tweedie link power. Defaults to 0.
theta Theta. Defaults to 0.
solver AUTO will set the solver based on given data and the other parameters. IRLSM is fast on on problems with small number of predictors and for lambda-search with L1 penalty, L_BFGS scales better for datasets with many columns. Must be one of: "AUTO", "IRLSM", "L_BFGS", "COORDINATE_DESCENT_NAIVE", "COORDINATE_DESCENT", "GRADIENT_DESCENT_LH", "GRADIENT_DESCENT_SQERR". Defaults to AUTO.
alpha Distribution of regularization between the L1 (Lasso) and L2 (Ridge) penalties. A value of 1 for alpha represents Lasso regression, a value of 0 produces Ridge regression, and anything in between specifies the amount of mixing between the two. Default value of alpha is 0 when SOLVER = 'L-BFGS'; 0.5 otherwise.
lambda Regularization strength.
lambda_search Logical. Use lambda search starting at lambda max, given lambda is then interpreted as lambda min. Defaults to FALSE.
early_stopping Logical. Stop early when there is no more relative improvement on train or validation (if provided). Defaults to TRUE.
nlambdas Number of lambdas to be used in a search. Default indicates: If alpha is zero, with lambda search set to True, the value of nlamdas is set to 30 (fewer lambdas are needed for ridge regression) otherwise it is set to 100. Defaults to -1.
**standardize**  Logical. Standardize numeric columns to have zero mean and unit variance. Defaults to FALSE.

**missing_values_handling**  Handling of missing values. Either MeanImputation, Skip or PlugValues. Must be one of: "MeanImputation", "Skip", "PlugValues". Defaults to MeanImputation.

**plug_values**  Plug Values (a single row frame containing values that will be used to impute missing values of the training/validation frame, use with conjunction missing_values_handling = PlugValues)

**compute_p_values**  Logical. Request p-values computation, p-values work only with IRLSM solver and no regularization. Defaults to FALSE.

**remove_collinear_columns**  Logical. In case of linearly dependent columns, remove some of the dependent columns. Defaults to FALSE.

**intercept**  Logical. Include constant term in the model. Defaults to TRUE.

**non_negative**  Logical. Restrict coefficients (not intercept) to be non-negative. Defaults to FALSE.

**max_iterations**  Maximum number of iterations. Defaults to -1.

**objective_epsilon**  Converge if objective value changes less than this. Default indicates: If lambda_search is set to True the value of objective_epsilon is set to .0001. If the lambda_search is set to False and lambda is equal to zero, the value of objective_epsilon is set to .000001, for any other value of lambda the default value of objective_epsilon is set to .0001. Defaults to -1.

**beta_epsilon**  Converge if beta changes less (using L-infinity norm) than beta epsilon. Only applies to IRLSM solver. Defaults to 0.0001.

**gradient_epsilon**  Converge if objective changes less (using L-infinity norm) than this, ONLY applies to L-BFGS solver. Default indicates: If lambda_search is set to False and lambda is equal to zero, the default value of gradient_epsilon is equal to .000001, otherwise the default value is .0001. If lambda_search is set to True, the conditional values above are 1E-8 and 1E-6 respectively. Defaults to -1.

**link**  Link function. Must be one of: "family_default", "identity", "logit", "log", "inverse", "tweedie", "ologit". Defaults to family_default.

**startval**  double array to initialize coefficients for GAM.

**prior**  Prior probability for y==1. To be used only for logistic regression. If the data has been sampled and the mean of response does not reflect reality. Defaults to -1.

**cold_start**  Logical. Only applicable to multiple alpha/lambdas when calling GLM from GAM. If false, build the next model for next set of alpha/lambdas starting from the values provided by current model. If true will start GLM model from scratch. Defaults to FALSE.

**lambda_min_ratio**  Minimum lambda used in lambda search, specified as a ratio of lambda_max (the smallest lambda that drives all coefficients to zero). Default indicates:
if the number of observations is greater than the number of variables, then
lambda_min_ratio is set to 0.0001; if the number of observations is less than
the number of variables, then lambda_min_ratio is set to 0.01. Defaults to -1.

beta_constraints
Beta constraints

max_active_predictors
Maximum number of active predictors during computation. Use as a stopping
criterion to prevent expensive model building with many predictors. Default
indicates: If the IRLSM solver is used, the value of max_active_predictors is set
to 5000 otherwise it is set to 10000000. Defaults to -1.

interactions
A list of predictor column indices to interact. All pairwise combinations will be
computed for the list.

interaction_pairs
A list of pairwise (first order) column interactions.

obj_reg
Likelihood divider in objective value computation, default is 1/nobs Defaults to
-1.

export_checkpoints_dir
Automatically export generated models to this directory.

stopping_rounds
Early stopping based on convergence of stopping_metric. Stop if simple moving
average of length k of the stopping_metric does not improve for k:=stopping_rounds
scoring events (0 to disable) Defaults to 0.

stopping_metric
Metric to use for early stopping (AUTO: logloss for classification, deviance
for regression and anomaly_score for Isolation Forest). Note that custom
and custom_increasing can only be used in GBM and DRF with the Python
client. Must be one of: "AUTO", "deviance", "logloss", "MSE", "RMSE",
"MAE", "RMSLE", "AUC", "AUCPR", "lift_top_group", "misclassification",
"mean_per_class_error", "custom", "custom_increasing". Defaults to AUTO.

stopping_tolerance
Relative tolerance for metric-based stopping criterion (stop if relative improve-
ment is not at least this much) Defaults to 0.001.

balance_classes
Logical. Balance training data class counts via over/under-sampling (for im-
balanced data). Defaults to FALSE.

class_sampling_factors
Desired over/under-sampling ratios per class (in lexicographic order). If not
specified, sampling factors will be automatically computed to obtain class bal-
ance during training. Requires balance_classes.

max_after_balance_size
Maximum relative size of the training data after balancing class counts (can be
less than 1.0). Requires balance_classes. Defaults to 5.0.

max_runtime_secs
Maximum allowed runtime in seconds for model training. Use 0 to disable.
Defaults to 0.

custom_metric_func
Reference to custom evaluation function, format: ‘language:keyName=funcName’
num_knots  Number of knots for gam predictors
knot_ids   String arrays storing frame keys of knots. One for each gam column set specified in gam_columns
standardize_tp_gam_cols Logical. standardize tp (thin plate) predictor columns Defaults to FALSE.
scale_tp_penalty_mat Logical. Scale penalty matrix for tp (thin plate) smoothers as in R Defaults to FALSE.
bs         Basis function type for each gam predictors, 0 for cr, 1 for thin plate regression with knots, 2 for thin plate regression with SVD. If specified, must be the same size as gam_columns
scale      Smoothing parameter for gam predictors. If specified, must be of the same length as gam_columns
keep_gam_cols Logical. Save keys of model matrix Defaults to FALSE.
auc_type   Set default multinomial AUC type. Must be one of: "AUTO", "NONE", "MACRO_OVR", "WEIGHTED_OVR", "MACRO_OVO", "WEIGHTED_OVO". Defaults to AUTO.

Examples

## Not run:
h2o.init()

# Run GAM of CAPSULE ~ AGE + RACE + PSA + DCAPS
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
prostate$CAPSULE <- as.factor(prostate$CAPSULE)
h2o.gam(y = "CAPSULE", x = c("RACE"), gam_columns = c("PSA"),
        training_frame = prostate, family = "binomial")

## End(Not run)

---

**h2o.gbm**  
Build gradient boosted classification or regression trees

**Description**

Builds gradient boosted classification trees and gradient boosted regression trees on a parsed data set. The default distribution function will guess the model type based on the response column type. In order to run properly, the response column must be an numeric for "gaussian" or an enum for "bernoulli" or "multinomial".
Usage

h2o.gbm(
    x,
    y,
    training_frame,
    model_id = NULL,
    validation_frame = NULL,
    nfolds = 0,
    keep_cross_validation_models = TRUE,
    keep_cross_validation_predictions = FALSE,
    keep_cross_validation_fold_assignment = FALSE,
    score_each_iteration = FALSE,
    score_tree_interval = 0,
    fold_assignment = c("AUTO", "Random", "Modulo", "Stratified"),
    fold_column = NULL,
    ignore_const_cols = TRUE,
    offset_column = NULL,
    weights_column = NULL,
    balance_classes = FALSE,
    class_sampling_factors = NULL,
    max_after_balance_size = 5,
    ntree = 50,
    max_depth = 5,
    min_rows = 10,
    nbins = 20,
    nbins_top_level = 1024,
    nbins_cats = 1024,
    r2_stopping = Inf,
    stopping_rounds = 0,
    stopping_metric = c("AUTO", "deviance", "logloss", "MSE", "RMSE", "MAE", "RMSLE",
                       "AUC", "AUCPR", "lift_top_group", "misclassification", "mean_per_class_error",
                       "custom", "custom_increasing"),
    stopping_tolerance = 0.001,
    max_runtime_secs = 0,
    seed = -1,
    build_tree_one_node = FALSE,
    learn_rate = 0.1,
    learn_rate_annealing = 1,
    distribution = c("AUTO", "bernoulli", "quasibinomial", "multinomial", "gaussian",
                    "poisson", "gamma", "tweedie", "laplace", "quantile", "huber", "custom"),
    quantile_alpha = 0.5,
    tweedie_power = 1.5,
    huber_alpha = 0.9,
    checkpoint = NULL,
    sample_rate = 1,
    sample_rate_per_class = NULL,
    col_sample_rate = 1,
    col_sample_rate_change_per_level = 1,
col_sample_rate_per_tree = 1,
min_split_improvement = 1e-05,
histogram_type = c("AUTO", "UniformAdaptive", "Random", "QuantilesGlobal",
   "RoundRobin"),
max_abs_leafnode_pred = Inf,
pred_noise_bandwidth = 0,
categorical_encoding = c("AUTO", "Enum", "OneHotInternal", "OneHotExplicit",
   "Binary", "Eigen", "LabelEncoder", "SortByResponse", "EnumLimited"),
calibrate_model = FALSE,
calibration_frame = NULL,
custom_metric_func = NULL,
custom_distribution_func = NULL,
export_checkpoints_dir = NULL,
monotone_constraints = NULL,
check_constant_response = TRUE,
gainslift_bins = -1,
auc_type = c("AUTO", "NONE", "MACRO_OVR", "WEIGHTED_OVR", "MACRO_OVO",
   "WEIGHTED_OVO"),
verbose = FALSE
)

Arguments

x (Optional) A vector containing the names or indices of the predictor variables to
use in building the model. If x is missing, then all columns except y are used.

y The name or column index of the response variable in the data. The response
must be either a numeric or a categorical/factor variable. If the response is
numeric, then a regression model will be trained, otherwise it will train a classi-
fication model.

training_frame Id of the training data frame.
model_id Destination id for this model; auto-generated if not specified.
validation_frame Id of the validation data frame.
nfolds Number of folds for K-fold cross-validation (0 to disable or >= 2). Defaults to
0.
keep_cross_validation_models Logical. Whether to keep the cross-validation models. Defaults to TRUE.
keep_cross_validation_predictions Logical. Whether to keep the predictions of the cross-validation models. De-
faults to FALSE.
keep_cross_validation_fold_assignment Logical. Whether to keep the cross-validation fold assignment. Defaults to FALSE.
score_each_iteration Logical. Whether to score during each iteration of model training. Defaults to FALSE.
score_tree_interval
Score the model after every so many trees. Disabled if set to 0. Defaults to 0.

fold_assignment
Cross-validation fold assignment scheme, if fold_column is not specified. The 'Stratified' option will stratify the folds based on the response variable, for classification problems. Must be one of: "AUTO", "Random", "Modulo", "Stratified". Defaults to AUTO.

fold_column
Column with cross-validation fold index assignment per observation.

ignore_const_cols
Logical. Ignore constant columns. Defaults to TRUE.

offset_column
Offset column. This will be added to the combination of columns before applying the link function.

weights_column
Column with observation weights. Giving some observation a weight of zero is equivalent to excluding it from the dataset; giving an observation a relative weight of 2 is equivalent to repeating that row twice. Negative weights are not allowed. Note: Weights are per-row observation weights and do not increase the size of the data frame. This is typically the number of times a row is repeated, but non-integer values are supported as well. During training, rows with higher weights matter more, due to the larger loss function pre-factor. If you set weight = 0 for a row, the returned prediction frame at that row is zero and this is incorrect. To get an accurate prediction, remove all rows with weight == 0.

balance_classes
Logical. Balance training data class counts via over/under-sampling (for imbalanced data). Defaults to FALSE.

class_sampling_factors
Desired over/under-sampling ratios per class (in lexicographic order). If not specified, sampling factors will be automatically computed to obtain class balance during training. Requires balance_classes.

max_after_balance_size
Maximum relative size of the training data after balancing class counts (can be less than 1.0). Requires balance_classes. Defaults to 5.0.

ntrees
Number of trees. Defaults to 50.

max_depth
Maximum tree depth (0 for unlimited). Defaults to 5.

min_rows
Fewest allowed (weighted) observations in a leaf. Defaults to 10.

nbins
For numerical columns (real/int), build a histogram of (at least) this many bins, then split at the best point Defaults to 20.

nbins_top_level
For numerical columns (real/int), build a histogram of (at most) this many bins at the root level, then decrease by factor of two per level Defaults to 1024.

nbins_cats
For categorical columns (factors), build a histogram of this many bins, then split at the best point. Higher values can lead to more overfitting. Defaults to 1024.

r2_stopping
r2_stopping is no longer supported and will be ignored if set - please use stopping_rounds, stopping_metric and stopping_tolerance instead. Previous version of H2O would stop making trees when the R^2 metric equals or exceeds this Defaults to 1.797693135e+308.
stopping_rounds
 Early stopping based on convergence of stopping_metric. Stop if simple moving average of length k of the stopping_metric does not improve for k:=stopping_rounds scoring events (0 to disable) Defaults to 0.

stopping_metric
 Metric to use for early stopping (AUTO: logloss for classification, deviance for regression and anomaly_score for Isolation Forest). Note that custom and custom_increasing can only be used in GBM and DRF with the Python client. Must be one of: "AUTO", "deviance", "logloss", "MSE", "RMSE", "MAE", "RMSLE", "AUC", "AUCPR", "lift_top_group", "misclassification", "mean_per_class_error", "custom", "custom_increasing". Defaults to AUTO.

stopping_tolerance
 Relative tolerance for metric-based stopping criterion (stop if relative improvement is not at least this much) Defaults to 0.001.

max_runtime_secs
 Maximum allowed runtime in seconds for model training. Use 0 to disable. Defaults to 0.

seed
 Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).

build_tree_one_node
 Logical. Run on one node only; no network overhead but fewer cpus used. Suitable for small datasets. Defaults to FALSE.

learn_rate
 Learning rate (from 0.0 to 1.0) Defaults to 0.1.

learn_rate_annealing
 Scale the learning rate by this factor after each tree (e.g., 0.99 or 0.999) Defaults to 1.

distribution
 Distribution function Must be one of: "AUTO", "bernoulli", "quasibinomial", "multinomial", "gaussian", "poisson", "gamma", "tweedie", "laplace", "quantile", "huber", "custom". Defaults to AUTO.

quantile_alpha
 Desired quantile for Quantile regression, must be between 0 and 1. Defaults to 0.5.

tweedie_power
 Tweedie power for Tweedie regression, must be between 1 and 2. Defaults to 1.5.

huber_alpha
 Desired quantile for Huber/M-regression (threshold between quadratic and linear loss, must be between 0 and 1). Defaults to 0.9.

checkpoint
 Model checkpoint to resume training with.

col_sample_rate
 Column sample rate (from 0.0 to 1.0) Defaults to 1.

col_sample_rate_change_per_level
 Relative change of the column sampling rate for every level (must be > 0.0 and <= 2.0) Defaults to 1.
col_sample_rate_per_tree
Column sample rate per tree (from 0.0 to 1.0) Defaults to 1.

min_split_improvement
Minimum relative improvement in squared error reduction for a split to happen
Defaults to 1e-05.

histogram_type
What type of histogram to use for finding optimal split points Must be one of:
"AUTO", "UniformAdaptive", "Random", "QuantilesGlobal", "RoundRobin". Defaults to AUTO.

max_abs_leafnode_pred
Maximum absolute value of a leaf node prediction Defaults to 1.797693135e+308.

pred_noise_bandwidth
Bandwidth (sigma) of Gaussian multiplicative noise ~N(1,sigma) for tree node
predictions Defaults to 0.

categorical_encoding
Encoding scheme for categorical features Must be one of: "AUTO", "Enum",
"OneHotInternal", "OneHotExplicit", "Binary", "Eigen", "LabelEncoder", "SortByResponse", "EnumLimited". Defaults to AUTO.

calibrate_model
Logical. Use Platt Scaling to calculate calibrated class probabilities. Calibration can provide more accurate estimates of class probabilities. Defaults to FALSE.

calibration_frame
Calibration frame for Platt Scaling

custom_metric_func
Reference to custom evaluation function, format: 'language:keyName=funcName'

custom_distribution_func
Reference to custom distribution, format: 'language:keyName=funcName'

export_checkpoints_dir
Automatically export generated models to this directory.

monotone_constraints
A mapping representing monotonic constraints. Use +1 to enforce an increasing constraint and -1 to specify a decreasing constraint.

check_constant_response
Logical. Check if response column is constant. If enabled, then an exception is thrown if the response column is a constant value. If disabled, then model will train regardless of the response column being a constant value or not. Defaults to TRUE.

gainslift_bins
Gains/Lift table number of bins. 0 means disabled. Default value -1 means automatic binning. Defaults to -1.

auc_type
Set default multinomial AUC type. Must be one of: "AUTO", "NONE", "MACRO_OVR",
"WEIGHTED_OVR", "MACRO_OVO", "WEIGHTED_OVO". Defaults to AUTO.

verbose
Logical. Print scoring history to the console (Metrics per tree). Defaults to FALSE.

See Also

predict.H2OModel for prediction
h2o.generic

Imports a generic model into H2O. Such model can be used then used for scoring and obtaining additional information about the model. The imported model has to be supported by H2O.

Description
Imports a generic model into H2O. Such model can be used then used for scoring and obtaining additional information about the model. The imported model has to be supported by H2O.

Usage

h2o.generic(model_id = NULL, model_key = NULL, path = NULL)

Arguments

- **model_id**: Destination id for this model; auto-generated if not specified.
- **model_key**: Key to the self-contained model archive already uploaded to H2O.
- **path**: Path to file with self-contained model archive.

Examples

```r
## Not run:
# library(h2o)
# h2o.init()

# generic_model <- h2o.genericModel(path="/path/to/model.zip", model_id="my_model")
# predictions <- h2o.predict(generic_model, dataset)
## End(Not run)
```
h2o.genericModel \ Imports a model under given path, creating a Generic model with it.

Description

Usage example: generic_model <- h2o.genericModel(model_file_path = "/path/to/mojo.zip") predictions <- h2o.predict(generic_model, dataset)

Usage

h2o.genericModel(mojo_file_path, model_id = NULL)

Arguments

mojo_file_path  Filesystem path to the model imported
model_id        Model ID, default is NULL

Value

Returns H2O Generic Model based on given embedded model

Examples

## Not run:

# Import default Iris dataset as H2O frame
data <- as.h2o(iris)

# Train a very simple GBM model
original_model <- h2o.gbm(x = features, y = "Species", training_frame = data)

# Download the trained GBM model as MOJO (temporary directory used in this example)
mojo_original_name <- h2o.download_mojo(model = original_model, path = tempdir())
mojo_original_path <- paste0(tempdir(), "/", mojo_original_name)

# Import the MOJO as Generic model
generic_model <- h2o.genericModel(mojo_original_path)

# Perform scoring with the generic model
generic_model_predictions <- h2o.predict(generic_model, data)

## End(Not run)
h2o.getAlphaBest  

**Description**

This function allows setting betas of an existing glm model.

**Usage**

```r
h2o.getAlphaBest(model)
```

**Arguments**

- `model`: an H2OModel corresponding from a h2o.glm call.

---

h2o.getConnection  

**Description**

Attempt to recover an h2o connection.

**Usage**

```r
h2o.getConnection()
```

**Value**

Returns an H2OConnection object.

---

h2o.getFrame  

**Description**

Get an R Reference to an H2O Dataset, that will NOT be GC'd by default

**Usage**

```r
h2o.getFrame(id)
```

**Arguments**

- `id`: A string indicating the unique frame of the dataset to retrieve.
Examples

```r
# Not run:
library(h2o)
h2o.init()

train <- h2o.importFile(f)
y <- "species"
x <- setdiff(names(train), y)
train[, y] <- as.factor(train[, y])
nfolds <- 5
num_base_models <- 2
my_gbm <- h2o.gbm(x = x, y = y, training_frame = train,
                   distribution = "multinomial", ntrees = 10,
                   max_depth = 3, min_rows = 2, learn_rate = 0.2,
                   nfolds = nfolds, fold_assignment = "Modulo",
                   keep_cross_validation_predictions = TRUE, seed = 1)
my_rf <- h2o.randomForest(x = x, y = y, training_frame = train,
                          ntree = 50, nfolds = nfolds, fold_assignment = "Modulo",
                          keep_cross_validation_predictions = TRUE, seed = 1)
stack <- h2o.stackedEnsemble(x = x, y = y, training_frame = train,
                          model_id = "my_ensemble_l1",
                          base_models = list(my_gbm@model_id, my_rf@model_id),
                          keep_levelone_frame = TRUE)
h2o.getFrame(stack@model$levelone_frame_id$name)
# End(Not run)
```

---

### h2o.getGLMFullRegularizationPath

**Extract full regularization path from a GLM model**

**Description**

Extract the full regularization path from a GLM model (assuming it was run with the lambda search option).

**Usage**

```r
h2o.getGLMFullRegularizationPath(model)
```

**Arguments**

- **model**: an `H2OModel` corresponding from a `h2o.glm` call.
h2o.getGrid

Get a grid object from H2O distributed K/V store.

Description

Note that if neither cross-validation nor a validation frame is used in the grid search, then the training metrics will display in the "get grid" output. If a validation frame is passed to the grid, and nfolds = 0, then the validation metrics will display. However, if nfolds > 1, then cross-validation metrics will display even if a validation frame is provided.

Usage

h2o.getGrid(grid_id, sort_by, decreasing, verbose = FALSE)

Arguments

grid_id ID of existing grid object to fetch
sort_by Sort the models in the grid space by a metric. Choices are "logloss", "residual_deviance", "mse", "auc", "accuracy", "precision", "recall", "f1", etc.
decreasing Specify whether sort order should be decreasing
verbose Controls verbosity of the output, if enabled prints out error messages for failed models (default: FALSE)

Examples

## Not run:
library(h2o)
library(jsonlite)
h2o.init()
iris_hf <- as.h2o(iris)
h2o.grid("gbm", grid_id = "gbm_grid_id", x = c(1:4), y = 5,
  training_frame = iris_hf, hyper_params = list(ntrees = c(1, 2, 3)))
grid <- h2o.getGrid("gbm_grid_id")
# Get grid summary
summary(grid)
# Fetch grid models
model_ids <- grid$model_ids
models <- lapply(model_ids, function(id) { h2o.getModel(id)})

## End(Not run)
h2o.getId

Get back-end distributed key/value store id from an H2OFrame.

Description
Get back-end distributed key/value store id from an H2OFrame.

Usage

h2o.getId(x)

Arguments

x
An H2OFrame

Value
The id of the H2OFrame

Examples

## Not run:
library(h2o)
h2o.init()

iris <- h2o.importFile(f)
h2o.getId(iris)

## End(Not run)

h2o.getLambdaBest

Extract best lambda value found from glm model.

Description
This function allows setting betas of an existing glm model.

Usage

h2o.getLambdaBest(model)

Arguments

model
an H2OModel corresponding from a h2o.glm call.
h2o.getLambdaMax  
Extract the maximum lambda value used during lambda search from glm model.

Description
This function allows setting betas of an existing glm model.

Usage
h2o.getLambdaMax(model)

Arguments
model  an H2OModel corresponding from a h2o.glm call.

h2o.getLambdaMin  
Extract the minimum lambda value calculated during lambda search from glm model. Note that due to early stop, this minimum lambda value may not be used in the actual lambda search.

Description
This function allows setting betas of an existing glm model.

Usage
h2o.getLambdaMin(model)

Arguments
model  an H2OModel corresponding from a h2o.glm call.
h2o.getModel

Get an R reference to an H2O model

Description

Returns a reference to an existing model in the H2O instance.

Usage

h2o.getModel(model_id)

Arguments

model_id

A string indicating the unique model_id of the model to retrieve.

Value

Returns an object that is a subclass of H2OModel.

Examples

## Not run:
library(h2o)
h2o.init()

iris_hf <- as.h2o(iris)
model_id <- h2o.gbm(x = 1:4, y = 5, training_frame = iris_hf)@model_id
model_retrieved <- h2o.getModel(model_id)

## End(Not run)

h2o.getModelTree

Fetches a single tree of a H2O model. This function is intended to be used on Gradient Boosting Machine models or Distributed Random Forest models.

Description

Fetches a single tree of a H2O model. This function is intended to be used on Gradient Boosting Machine models or Distributed Random Forest models.

Usage

h2o.getModelTree(
  model,
  tree_number,
  tree_class = NA,
  plain_language_rules = "AUTO"
)
Arguments

- **model**: Model with trees
- **tree_number**: Number of the tree in the model to fetch, starting with 1
- **tree_class**: Name of the class of the tree (if applicable). This value is ignored for regression and binomial response column, as there is only one tree built. As there is exactly one class per categorical level, name of tree’s class equals to the corresponding categorical level of response column.
- **plain_language_rules**: (Optional) Whether to generate plain language rules. AUTO by default, meaning FALSE for big trees and TRUE for small trees.

Value

Returns an H2OTree object with detailed information about a tree.

Examples

```r
## Not run:
library(h2o)
h2o.init()

iris <- h2o.importFile(f)
gbm_model <- h2o.gbm(y = "species", training_frame = iris)
tree <- h2o.getModelTree(gbm_model, 1, "Iris-setosa")

## End(Not run)
```

---

**h2o.getTimezone**

*Get the Time Zone on the H2O cluster Returns a string*

**Description**

Get the Time Zone on the H2O cluster Returns a string

**Usage**

```r
h2o.getTimezone()
```
**h2o.getTypes**

*Get the types-per-column*

**Description**

Get the types-per-column

**Usage**

```r
h2o.getTypes(x)
```

**Arguments**

- `x` An H2OFrame

**Value**

A list of types per column

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

iris <- h2o.importFile(f)
h2o.getTypes(iris)

## End(Not run)
```

**h2o.getVersion**

*Get h2o version*

**Description**

Get h2o version

**Usage**

```r
h2o.getVersion()
```
h2o.get_automl

Get an R object that is a subclass of H2OAutoML

Description

Get an R object that is a subclass of H2OAutoML.

Usage

h2o.get_automl(project_name)

h2o.getAutoML(project_name)

Arguments

project_name A string indicating the project_name of the automl instance to retrieve.

Value

Returns an object that is a subclass of H2OAutoML.

Examples

## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.importFile(path = prostate_path, header = TRUE)
y <- "CAPSULE"
prostate[,y] <- as.factor(prostate[,y]) #convert to factor for classification
aml <- h2o.automl(y = y, training_frame = prostate,
                   max_runtime_secs = 30, project_name = "prostate")
aml2 <- h2o.get_automl("prostate")

## End(Not run)
Usage

```r
h2o.get_best_model(
  object,
  algorithm = c("any", "basemodel", "deeplearning", "drf", "gbm", "glm",
                "stackedensemble", "xgboost"),
  criterion = c("AUTO", "AUC", "AUCPR", "logloss", "MAE", "mean_per_class_error",
               "deviance", "MSE", "predict_time_per_row_ms", "RMSE", "RMSLE", "training_time_ms")
)
```

Arguments

- **object**: H2OAutoML object
- **algorithm**: One of "any", "basemodel", "deeplearning", "drf", "gbm", "glm", "stackedensemble", "xgboost"
- **criterion**: Criterion can be one of the metrics reported in the leaderboard. If set to NULL, the same ordering as in the leaderboard will be used. Available criteria:
  - Regression metrics: deviance, RMSE, MSE, MAE, RMSLE
  - Binomial metrics: AUC, logloss, AUCPR, mean_per_class_error, RMSE, MSE
  - Multinomial metrics: mean_per_class_error, logloss, RMSE, MSE

The following additional leaderboard information can be also used as a criterion:
- 'training_time_ms': column providing the training time of each model in milliseconds (doesn't include the training of cross validation models).
- 'predict_time_per_row_ms': column providing the average prediction time by the model for a single row.

Value

An H2OModel or NULL if no model of a given family is present

Examples

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.importFile(path = prostate_path, header = TRUE)
y <- "CAPSULE"
prostate[,y] <- as.factor(prostate[,y]) #convert to factor for classification
aml <- h2o.automl(y = y, training_frame = prostate, max_runtime_secs = 30)
gbm <- h2o.get_best_model(aml, "gbm")
## End(Not run)
```
h2o.get_best_model_predictors

Extracts the subset of predictor names that yield the best R2 value for each predictor subset size.

Description

Extracts the subset of predictor names that yield the best R2 value for each predictor subset size.

Usage

h2o.get_best_model_predictors(model)

Arguments

model is a H2OModel with algorithm name of modelselection

h2o.get_best_r2_values

Extracts the best R2 values for all predictor subset size.

Description

Extracts the best R2 values for all predictor subset size.

Usage

h2o.get_best_r2_values(model)

Arguments

model is a H2OModel with algorithm name of modelselection
h2o.get_leaderboard

Retrieve the leaderboard from the AutoML instance.

Description

Contrary to the default leaderboard attached to the automl instance, this one can return columns other than the metrics.

Usage

h2o.get_leaderboard(object, extra_columns = NULL)

Arguments

object

The object for which to return the leaderboard. Currently, only H2OAutoML instances are supported.

extra_columns

A string or a list of string specifying which optional columns should be added to the leaderboard. Defaults to None. Currently supported extensions are:

• 'ALL': adds all columns below.
• 'training_time_ms': column providing the training time of each model in milliseconds (doesn’t include the training of cross validation models).
• 'predict_time_per_row_ms': column providing the average prediction time by the model for a single row.
• 'algo': column providing the algorithm name for each model.

Value

An H2OFrame representing the leaderboard.

Examples

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.importFile(path = prostate_path, header = TRUE)
y <- "CAPSULE"
prostate[,y] <- as.factor(prostate[,y])  #convert to factor for classification
aml <- h2o.automl(y = y, training_frame = prostate, max_runtime_secs = 30)
lb <- h2o.get_leaderboard(aml)
head(lb)

## End(Not run)
```
h2o.get_segment_models

Retrieves an instance of `H2OSegmentModels` for a given id.

Description

Retrieves an instance of `H2OSegmentModels` for a given id.

Usage

```r
h2o.get_segment_models(segment_models_id)
```

Arguments

- `segment_models_id`:
  A string indicating the unique `segment_models_id`

Value

Returns an object that is a subclass of `H2OSegmentModels`.

Examples

```r
## Not run:
library(h2o)
h2o.init()
iris_hf <- as.h2o(iris)
h2o.train_segments(algorithm = "gbm",
                   segment_columns = "Species",
                   segment_models_id="models_by_species",
                   x = c(1:3), y = 4, training_frame = iris_hf, ntrees = 5, max_depth = 4)
models <- h2o.get_segment_models("models_by_species")
as.data.frame(models)

## End(Not run)
```
h2o.giniCoef | Retrieve the GINI Coefficient

Description

Retrieves the GINI coefficient from an H2OBinomialMetrics. If "train", "valid", and "xval" parameters are FALSE (default), then the training GINI value is returned. If more than one parameter is set to TRUE, then a named vector of GINIs are returned, where the names are "train", "valid" or "xval".

Usage

h2o.giniCoef(object, train = FALSE, valid = FALSE, xval = FALSE)

Arguments

object | an H2OBinomialMetrics object.
train | Retrieve the training GINI Coefficient
valid | Retrieve the validation GINI Coefficient
xval | Retrieve the cross-validation GINI Coefficient

See Also

h2o.auc for AUC, h2o.giniCoef for the GINI coefficient, and h2o.metric for the various threshold metrics. See h2o.performance for creating H2OModelMetrics objects.

Examples

```r
## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(prostate_path)
prostate[, 2] <- as.factor(prostate[, 2])
model <- h2o.gbm(x = 3:9, y = 2, training_frame = prostate, distribution = "bernoulli")
perf <- h2o.performance(model, prostate)
h2o.giniCoef(perf)

## End(Not run)
```
**h2o.glm**

*Fit a generalized linear model*

**Description**

Fits a generalized linear model, specified by a response variable, a set of predictors, and a description of the error distribution.

**Usage**

```r
h2o.glm(
  x,
  y,
  training_frame,
  model_id = NULL,
  validation_frame = NULL,
  nfolds = 0,
  checkpoint = NULL,
  export_checkpoints_dir = NULL,
  seed = -1,
  keep_cross_validation_models = TRUE,
  keep_cross_validation_predictions = FALSE,
  keep_cross_validation_fold_assignment = FALSE,
  fold_assignment = c("AUTO", "Random", "Modulo", "Stratified"),
  fold_column = NULL,
  random_columns = NULL,
  ignore_const_cols = TRUE,
  score_each_iteration = FALSE,
  score_iteration_interval = -1,
  offset_column = NULL,
  weights_column = NULL,
  family = c("AUTO", "gaussian", "binomial", "fractionalbinomial", "quasibinomial",
             "ordinal", "multinomial", "poisson", "gamma", "tweedie", "negativebinomial"),
  rand_family = c("[gaussian]"),
  tweedie_variance_power = 0,
  tweedie_link_power = 1,
  theta = 1e-10,
  solver = c("AUTO", "IRLSM", "L_BFGS", "COORDINATE_DESCENT_NAIVE",
             "COORDINATE_DESCENT", "GRADIENT_DESCENT_LH", "GRADIENT_DESCENT_SQERR"),
  alpha = NULL,
  lambda = NULL,
  lambda_search = FALSE,
  early_stopping = TRUE,
  nlambdas = -1,
  standardize = TRUE,
  missing_values_handling = c("MeanImputation", "Skip", "PlugValues"),
  plug_values = NULL,
)```

compute_p_values = FALSE,
remove_collinear_columns = FALSE,
intercept = TRUE,
non_negative = FALSE,
max_iterations = -1,
objective_epsilon = -1,
beta_epsilon = 1e-04,
gradient_epsilon = -1,
link = c("family_default", "identity", "logit", "log", "inverse", "tweedie",
          "ologit"),
rand_link = c("[identity]", "[family_default]"),
startval = NULL,
calc_like = FALSE,
HGLM = FALSE,
prior = -1,
cold_start = FALSE,
lambda_min_ratio = -1,
beta_constraints = NULL,
max_active_predictors = -1,
interactions = NULL,
interaction_pairs = NULL,
obj_reg = -1,
stopping_rounds = 0,
stopping_metric = c("AUTO", "deviance", "logloss", "MSE", "RMSE", "MAE", "RMSLE",
                    "AUC", "AUCPR", "lift_top_group", "misclassification", "mean_per_class_error",
                    "custom", "custom_increasing"),
stopping_tolerance = 0.001,
balance_classes = FALSE,
class_sampling_factors = NULL,
max_after_balance_size = 5,
max_runtime_secs = 0,
custom_metric_func = NULL,
generate_scoring_history = FALSE,
auc_type = c("AUTO", "NONE", "MACRO_OVR", "WEIGHTED_OVR", "MACRO_OVO",
            "WEIGHTED_OVO")
)

Arguments

x  (Optional) A vector containing the names or indices of the predictor variables to use in building the model. If x is missing, then all columns except y are used.

y  The name or column index of the response variable in the data. The response must be either a numeric or a categorical/factor variable. If the response is numeric, then a regression model will be trained, otherwise it will train a classification model.

training_frame  Id of the training data frame.

model_id  Destination id for this model; auto-generated if not specified.
validation_frame
  Id of the validation data frame.

nfolds
  Number of folds for K-fold cross-validation (0 to disable or >= 2). Defaults to 0.

checkpoint
  Model checkpoint to resume training with.

export_checkpoints_dir
  Automatically export generated models to this directory.

seed
  Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).

keep_cross_validation_models
  Logical. Whether to keep the cross-validation models. Defaults to TRUE.

keep_cross_validation_predictions
  Logical. Whether to keep the predictions of the cross-validation models. Defaults to FALSE.

keep_cross_validation_fold_assignment
  Logical. Whether to keep the cross-validation fold assignment. Defaults to FALSE.

fold_assignment
  Cross-validation fold assignment scheme, if fold_column is not specified. The 'Stratified' option will stratify the folds based on the response variable, for classification problems. Must be one of: "AUTO", "Random", "Modulo", "Stratified". Defaults to AUTO.

fold_column
  Column with cross-validation fold index assignment per observation.

random_columns
  random columns indices for HGLM.

ignore_const_cols
  Logical. Ignore constant columns. Defaults to TRUE.

score_each_iteration
  Logical. Whether to score during each iteration of model training. Defaults to FALSE.

score_iteration_interval
  Perform scoring for every score_iteration_interval iterations Defaults to -1.

offset_column
  Offset column. This will be added to the combination of columns before applying the link function.

weights_column
  Column with observation weights. Giving some observation a weight of zero is equivalent to excluding it from the dataset; giving an observation a relative weight of 2 is equivalent to repeating that row twice. Negative weights are not allowed. Note: Weights are per-row observation weights and do not increase the size of the data frame. This is typically the number of times a row is repeated, but non-integer values are supported as well. During training, rows with higher weights matter more, due to the larger loss function pre-factor. If you set weight = 0 for a row, the returned prediction frame at that row is zero and this is incorrect. To get an accurate prediction, remove all rows with weight == 0.

family
  Family. Use binomial for classification with logistic regression, others are for regression problems. Must be one of: "AUTO", "gaussian", "binomial", "fractionalbinomial", "quasibinomial", "ordinal", "multinomial", "poisson", "gamma", "tweedie", "negativebinomial". Defaults to AUTO.
**rand_family**  Random Component Family array. One for each random component. Only support gaussian for now. Must be one of: "[gaussian]".

**tweedie_variance_power**  Tweedie variance power Defaults to 0.

**tweedie_link_power**  Tweedie link power Defaults to 1.

**theta**  Theta Defaults to 1e-10.

**solver**  AUTO will set the solver based on given data and the other parameters. IRLSM is fast on on problems with small number of predictors and for lambda-search with L1 penalty, _L_BFGS_ scales better for datasets with many columns. Must be one of: "AUTO", "IRLSM", "L_BFGS", "COORDINATE_DESCENT_NAIVE", "COORDINATE_DESCENT", "GRADIENT_DESCENT_LH", "GRADIENT_DESCENT_SQERR". Defaults to AUTO.

**alpha**  Distribution of regularization between the L1 (Lasso) and L2 (Ridge) penalties. A value of 1 for alpha represents Lasso regression, a value of 0 produces Ridge regression, and anything in between specifies the amount of mixing between the two. Default value of alpha is 0 when SOLVER = 'L-BFGS'; 0.5 otherwise.

**lambda**  Regularization strength

**lambda_search**  Logical. Use lambda search starting at lambda max, given lambda is then interpreted as lambda min Defaults to FALSE.

**early_stopping**  Logical. Stop early when there is no more relative improvement on train or validation (if provided) Defaults to TRUE.

**nlambdas**  Number of lambdas to be used in a search. Default indicates: If alpha is zero, with lambda search set to True, the value of nlamdas is set to 30 (fewer lambdas are needed for ridge regression) otherwise it is set to 100. Defaults to -1.

**standardize**  Logical. Standardize numeric columns to have zero mean and unit variance Defaults to TRUE.

**missing_values_handling**  Handling of missing values. Either MeanImputation, Skip or PlugValues. Must be one of: "MeanImputation", "Skip", "PlugValues". Defaults to MeanImputation.

**plug_values**  Plug Values (a single row frame containing values that will be used to impute missing values of the training/validation frame, use with conjunction missing_values_handling = PlugValues)

**compute_p_values**  Logical. Request p-values computation, p-values work only with IRLSM solver and no regularization Defaults to FALSE.

**remove_collinear_columns**  Logical. In case of linearly dependent columns, remove some of the dependent columns Defaults to FALSE.

**intercept**  Logical. Include constant term in the model Defaults to TRUE.

**non_negative**  Logical. Restrict coefficients (not intercept) to be non-negative Defaults to FALSE.

**max_iterations**  Maximum number of iterations Defaults to -1.
objective_epsilon
Converge if objective value changes less than this. Default indicates: If lambda_search
is set to True the value of objective_epsilon is set to .0001. If the lambda_search
is set to False and lambda is equal to zero, the value of objective_epsilon is set
to .000001, for any other value of lambda the default value of objective_epsilon
is set to .0001. Defaults to -1.

beta_epsilon
Converge if beta changes less (using L-infinity norm) than beta epsilon, ONLY
applies to IRLSM solver. Defaults to 0.0001.

gradient_epsilon
Converge if objective changes less (using L-infinity norm) than this, ONLY ap-
plies to L-BFGS solver. Default indicates: If lambda_search is set to False
and lambda is equal to zero, the default value of gradient_epsilon is equal to
.000001, otherwise the default value is .0001. If lambda_search is set to True,
the conditional values above are 1E-8 and 1E-6 respectively. Defaults to -1.

link
Link function. Must be one of: "family_default", "identity", "logit", "log", "in-
verse", "tweedie", "ologit". Defaults to family_default.

rand_link
Link function array for random component in HGLM. Must be one of: ":[identity], ":[family_default]"

startval
double array to initialize fixed and random coefficients for HGLM, coefficients
for GLM.

calc_like
Logical. if true, will return likelihood function value for HGLM. Defaults to
FALSE.

HGLM
Logical. If set to true, will return HGLM model. Otherwise, normal GLM
model will be returned. Defaults to FALSE.

prior
Prior probability for y==1. To be used only for logistic regression iff the data
has been sampled and the mean of response does not reflect reality. Defaults to
-1.

cold_start
Logical. Only applicable to multiple alpha/lambda values. If false, build the
next model for next set of alpha/lambda values starting from the values pro-
vided by current model. If true will start GLM model from scratch. Defaults to
FALSE.

lambda_min_ratio
Minimum lambda used in lambda search, specified as a ratio of lambda_max
(the smallest lambda that drives all coefficients to zero). Default indicates:
if the number of observations is greater than the number of variables, then
lambda_min_ratio is set to 0.0001; if the number of observations is less than
the number of variables, then lambda_min_ratio is set to 0.01. Defaults to -1.

beta_constraints
Beta constraints

max_active_predictors
Maximum number of active predictors during computation. Use as a stopping
criterion to prevent expensive model building with many predictors. Default
indicates: If the IRLSM solver is used, the value of max_active_predictors is set
to 5000 otherwise it is set to 100000000. Defaults to -1.

interactions
A list of predictor column indices to interact. All pairwise combinations will be
computed for the list.
interaction_pairs
A list of pairwise (first order) column interactions.

obj_reg
Likelihood divider in objective value computation, default is 1/nobs Defaults to 1.

stopping_rounds
Early stopping based on convergence of stopping_metric. Stop if simple moving average of length k of the stopping_metric does not improve for k:=stopping_rounds scoring events (0 to disable) Defaults to 0.

stopping_metric
Metric to use for early stopping (AUTO: logloss for classification, deviance for regression and anomaly_score for Isolation Forest). Note that custom and custom_increasing can only be used in GBM and DRF with the Python client. Must be one of: "AUTO", "deviance", "logloss", "MSE", "RMSE", "MAE", "RMSLE", "AUC", "AUCPR", "lift_top_group", "misclassification", "mean_per_class_error", "custom", "custom_increasing". Defaults to AUTO.

stopping_tolerance
Relative tolerance for metric-based stopping criterion (stop if relative improvement is not at least this much) Defaults to 0.001.

balance_classes
Logical. Balance training data class counts via over/under-sampling (for imbalanced data). Defaults to FALSE.

class_sampling_factors
Desired over/under-sampling ratios per class (in lexicographic order). If not specified, sampling factors will be automatically computed to obtain class balance during training. Requires balance_classes.

max_after_balance_size
Maximum relative size of the training data after balancing class counts (can be less than 1.0). Requires balance_classes. Defaults to 5.0.

max_runtime_secs
Maximum allowed runtime in seconds for model training. Use 0 to disable. Defaults to 0.

custom_metric_func
Reference to custom evaluation function, format: ‘language:keyName=funcName’

generate_scoring_history
Logical. If set to true, will generate scoring history for GLM. This may significantly slow down the algo. Defaults to FALSE.

auc_type
Set default multinomial AUC type. Must be one of: "AUTO", "NONE", "MACRO_OVR", "WEIGHTED_OVR", "MACRO_OVO", "WEIGHTED_OVO". Defaults to AUTO.

Value
A subclass of H2OModel is returned. The specific subclass depends on the machine learning task at hand (if it's binomial classification, then an H2OBinomialModel is returned, if it's regression then a H2ORegressionModel is returned). The default print-out of the models is shown, but further GLM-specific information can be queried out of the object. To access these various items, please refer to the seealso section below. Upon completion of the GLM, the resulting object has coefficients, normalized coefficients, residual/null deviance, aic, and a host of model metrics including
MSE, AUC (for logistic regression), degrees of freedom, and confusion matrices. Please refer to the more in-depth GLM documentation available here: https://h2o-release.s3.amazonaws.com/h2o-dev/rel-shannon/2/docs-website/h2o-docs/index.html#Data+Science+Algorithms-GLM

See Also
predict.H2OModel for prediction, h2o.mse, h2o.auc, h2o.confusionMatrix, h2o.performance, h2o.giniCoef, h2o.logloss, h2o.varimp, h2o.scoreHistory

Examples

```r
## Not run:
h2o.init()

# Run GLM of CAPSULE ~ AGE + RACE + PSA + DCAPS
prostate_path = system.file("extdata", "prostate.csv", package = "h2o")
prostate = h2o.importFile(path = prostate_path)
h2o.glm(y = "CAPSULE", x = c("AGE", "RACE", "PSA", "DCAPS"), training_frame = prostate,
       family = "binomial", nfolds = 0, alpha = 0.5, lambda_search = FALSE)

# Run GLM of VOL ~ CAPSULE + AGE + RACE + PSA + GLEASON
predictors = setdiff(colnames(prostate), c("ID", "DPROS", "DCAPS", "VOL"))
h2o.glm(y = "VOL", x = predictors, training_frame = prostate, family = "gaussian",
        nfolds = 0, alpha = 0.1, lambda_search = FALSE)

# GLM variable importance
# Also see:
# https://github.com/h2oai/h2o/blob/master/R/tests/tdemos/runit_demo_VI_all_algos.R
bank = h2o.importFile(path="https://s3.amazonaws.com/h2o-public-test-data/smalldata/demos/bank-additional-full.csv")
predictors = 1:20
target = "y"
glm = h2o.glm(x = predictors,
      y = target,
      training_frame = bank,
      family = "binomial",
      standardize = TRUE,
      lambda_search = TRUE)
h2o.std_coef_plot(glm, num_of_features = 20)
## End(Not run)
```

h2o.glm

---

**Generalized low rank decomposition of an H2O data frame**

**Description**

Builds a generalized low rank decomposition of an H2O data frame.
Usage

h2o.glrm(
    training_frame,
    cols = NULL,
    model_id = NULL,
    validation_frame = NULL,
    ignore_const_cols = TRUE,
    score_each_iteration = FALSE,
    representation_name = NULL,
    loading_name = NULL,
    transform = c("NONE", "STANDARDIZE", "NORMALIZE", "DEMEAN", "DESCALE"),
    k = 1,
    loss = c("Quadratic", "Absolute", "Huber", "Poisson", "Hinge", "Logistic",
             "Periodic"),
    loss_by_col = c("Quadratic", "Absolute", "Huber", "Poisson", "Hinge", "Logistic",
                   "Periodic", "Categorical", "Ordinal"),
    loss_by_col_idx = NULL,
    multi_loss = c("Categorical", "Ordinal"),
    period = 1,
    regularization_x = c("None", "Quadratic", "L2", "L1", "NonNegative", "OneSparse",
                         "UnitOneSparse", "Simplex"),
    regularization_y = c("None", "Quadratic", "L2", "L1", "NonNegative", "OneSparse",
                         "UnitOneSparse", "Simplex"),
    gamma_x = 0,
    gamma_y = 0,
    max_iterations = 1000,
    max_updates = 2000,
    init_step_size = 1,
    min_step_size = 1e-04,
    seed = -1,
    init = c("Random", "SVD", "PlusPlus", "User"),
    svd_method = c("GramSVD", "Power", "Randomized"),
    user_y = NULL,
    user_x = NULL,
    expand_user_y = TRUE,
    impute_original = FALSE,
    recover_svd = FALSE,
    max_runtime_secs = 0,
    export_checkpoints_dir = NULL
)

Arguments

training_frame  Id of the training data frame.
cols           (Optional) A vector containing the data columns on which k-means operates.
model_id       Destination id for this model; auto-generated if not specified.
validation_frame  Id of the validation data frame.
ignore_const_cols
  Logical. Ignore constant columns. Defaults to TRUE.

score_each_iteration
  Logical. Whether to score during each iteration of model training. Defaults to FALSE.

representation_name
  Frame key to save resulting X

loading_name
  [Deprecated] Use representation_name instead. Frame key to save resulting X.

transform
  Transformation of training data Must be one of: "NONE", "STANDARDIZE", "NORMALIZE", "DEMEAN", "DESCALE". Defaults to NONE.

k
  Rank of matrix approximation Defaults to 1.

loss

loss_by_col
  Loss function by column (override) Must be one of: "Quadratic", "Absolute", "Huber", "Poisson", "Hinge", "Logistic", "Periodic", "Categorical", "Ordinal".

loss_by_col_idx
  Loss function by column index (override)

multi_loss
  Categorical loss function Must be one of: "Categorical", "Ordinal". Defaults to Categorical.

period
  Length of period (only used with periodic loss function) Defaults to 1.

regularization_x
  Regularization function for X matrix Must be one of: "None", "Quadratic", "L2", "L1", "NonNegative", "OneSparse", "UnitOneSparse", "Simplex". Defaults to None.

regularization_y
  Regularization function for Y matrix Must be one of: "None", "Quadratic", "L2", "L1", "NonNegative", "OneSparse", "UnitOneSparse", "Simplex". Defaults to None.

gamma_x
  Regularization weight on X matrix Defaults to 0.

gamma_y
  Regularization weight on Y matrix Defaults to 0.

max_iterations
  Maximum number of iterations Defaults to 1000.

max_updates
  Maximum number of updates, defaults to 2*max_iterations Defaults to 2000.

init_step_size
  Initial step size Defaults to 1.

min_step_size
  Minimum step size Defaults to 0.0001.

seed
  Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).

init
  Initialization mode Must be one of: "Random", "SVD", "PlusPlus", "User". Defaults to PlusPlus.

svd_method
  Method for computing SVD during initialization (Caution: Randomized is currently experimental and unstable) Must be one of: "GramSVD", "Power", "Randomized". Defaults to Randomized.

user_y
  User-specified initial Y
user_x User-specified initial X
expand_user_y Logical. Expand categorical columns in user-specified initial Y Defaults to TRUE.
impute_original Logical. Reconstruct original training data by reversing transform Defaults to FALSE.
recover_svd Logical. Recover singular values and eigenvectors of XY Defaults to FALSE.
max_runtime_secs Maximum allowed runtime in seconds for model training. Use 0 to disable. Defaults to 0.
export_checkpoints_dir Automatically export generated models to this directory.

Value

an object of class H2ODimReductionModel.

References


See Also

h2o.kmeans,h2o.svd,h2o.pcomp

Examples

## Not run:
library(h2o)
h2o.init()
australia_path <- system.file("extdata", "australia.csv", package = "h2o")
australia <- h2o.uploadFile(path = australia_path)
h2o.glrm(training_frame = australia, k = 5, loss = "Quadratic", regularization_x = "L1",
          gamma_x = 0.5, gamma_y = 0, max_iterations = 1000)

## End(Not run)
h2o.grep  

**Search for matches to an argument pattern**

**Description**

Searches for matches to argument ‘pattern’ within each element of a string column.

**Usage**

```r
h2o.grep(
  pattern,
  x,
  ignore.case = FALSE,
  invert = FALSE,
  output.logical = FALSE
)
```

**Arguments**

- `pattern`  
  A character string containing a regular expression.
- `x`  
  An H2O frame that wraps a single string column.
- `ignore.case`  
  If TRUE case is ignored during matching.
- `invert`  
  Identify elements that do not match the pattern.
- `output.logical`  
  If TRUE returns logical vector of indicators instead of list of matching positions

**Details**

This function has similar semantics as R’s native grep function and it supports a subset of its parameters. Default behavior is to return indices of the elements matching the pattern. Parameter ‘output.logical’ can be used to return a logical vector indicating if the element matches the pattern (1) or not (0).

**Value**

H2OFrame holding the matching positions or a logical vector if ‘output.logical’ is enabled.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
addresses <- as.h2o(c("2307", "Leghorn St", "Mountain View", "CA", "94043"))
zip_codes <- addresses[h2o.grep("[0-9]{5}", addresses, output.logical = TRUE),]
## End(Not run)
```
Description

Provides a set of functions to launch a grid search and get its results.

Usage

\[
\text{h2o.grid(}
\begin{align*}
\text{algorithm}, \\
\text{grid_id,} \\
\text{x,} \\
\text{y,} \\
\text{training_frame,} \\
\text{...}, \\
\text{hyper_params = list(),} \\
\text{is_supervised = NULL,} \\
\text{do_hyper_params_check = FALSE,} \\
\text{search_criteria = NULL,} \\
\text{export_checkpoints_dir = NULL,} \\
\text{recovery_dir = NULL,} \\
\text{parallelism = 1}
\end{align*}
\]

Arguments

- **algorithm**: Name of algorithm to use in grid search (gbm, randomForest, kmeans, glm, deeplearning, naivebayes, pca).
- **grid_id**: (Optional) ID for resulting grid search. If it is not specified then it is autogenerated.
- **x**: (Optional) A vector containing the names or indices of the predictor variables to use in building the model. If x is missing, then all columns except y are used.
- **y**: The name or column index of the response variable in the data. The response must be either a numeric or a categorical/factor variable. If the response is numeric, then a regression model will be trained, otherwise it will train a classification model.
- **training_frame**: Id of the training data frame.
- **hyper_params**: List of lists of hyper parameters (i.e., list(ntrees=c(1,2),max_depth=c(5,7))).
- **is_supervised**: [Deprecated] It is not possible to override default behaviour. (Optional) If specified then override the default heuristic which decides if the given algorithm name and parameters specify a supervised or unsupervised algorithm.
do_hyper_params_check
Perform client check for specified hyper parameters. It can be time expensive for large hyper space.

search_criteria
(Optional) List of control parameters for smarter hyperparameter search. The list can include values for: strategy, max_models, max_runtime_secs, stopping_metric, stopping_tolerance, stopping_rounds and seed. The default strategy 'Cartesian' covers the entire space of hyperparameter combinations. If you want to use cartesian grid search, you can leave the search_criteria argument unspecified. Specify the "RandomDiscrete" strategy to get random search of all the combinations of your hyperparameters with three ways of specifying when to stop the search: max number of models, max time, and metric-based early stopping (e.g., stop if MSE has not improved by 0.0001 over the 5 best models). Examples below: list(strategy = "RandomDiscrete",max_runtime_secs = 600,max_models = 100, stopping_metric = "AUTO", stopping_tolerance = 0.00001, stopping_rounds = 5, seed = 123456) or list(strategy = "RandomDiscrete",max_models = 42,max_runtime_secs = 28800) or list(strategy = "RandomDiscrete",stopping_metric = "AUTO",stopping_tolerance = 0.001, stopping_rounds = 10) or list(strategy = "RandomDiscrete",stopping_metric = "misclassification",stopping_tolerance = 0.00001, stopping_rounds = 5).

export_checkpoints_dir
Directory to automatically export grid and its models to.

recovery_dir
When specified the grid and all necessary data (frames, models) will be saved to this directory (use HDFS or other distributed file-system). Should the cluster crash during training, the grid can be reloaded from this directory via h2o.loadGrid and training can be resumed.

parallelism
Level of Parallelism during grid model building. 1 = sequential building (default). Use the value of 0 for adaptive parallelism - decided by H2O. Any number > 1 sets the exact number of models built in parallel.

Details
Launch grid search with given algorithm and parameters.

Examples
```r
## Not run:
library(h2o)
library(jsonlite)
h2o.init()
iris_hf <- as.h2o(iris)
grid <- h2o.grid("gbm", x = c(1:4), y = 5, training_frame = iris_hf,
                   hyper_params = list(ntrees = c(1, 2, 3)))
# Get grid summary
summary(grid)
# Fetch grid models
model_ids <- grid$model_ids
models <- lapply(model_ids, function(id) { h2o.getModel(id) })
```

## End(Not run)
h2o.group_by

Group and Apply by Column

Description

Performs a group by and apply similar to dply.

Usage

h2o.group_by(
  data, 
  by, 
  ..., 
  gb.control = list(na.methods = NULL, col.names = NULL)
)

Arguments

data: an H2OFrame object.
by: a list of column names
...: any supported aggregate function. See Details: for more help.
gb.control: a list of how to handle NA values in the dataset as well as how to name output columns. The method is specified using the rm.method argument. See Details: for more help.

Details

In the case of na.methods within gb.control, there are three possible settings. "all" will include NAs in computation of functions. "rm" will completely remove all NA fields. "ignore" will remove NAs from the numerator but keep the rows for computational purposes. If a list smaller than the number of columns groups is supplied, the list will be padded by "ignore".

Note that to specify a list of column names in the gb.control list, you must add the col.names argument. Similar to na.methods, col.names will pad the list with the default column names if the length is less than the number of columns groups supplied.

Supported functions include nrow. This function is required and accepts a string for the name of the generated column. Other supported aggregate functions accept col and na arguments for specifying columns and the handling of NAs ("all", "ignore", and GroupBy object; max calculates the maximum of each column specified in col for each group of a GroupBy object; mean calculates the mean of each column specified in col for each group of a GroupBy object; min calculates the minimum of each column specified in col for each group of a GroupBy object; node calculates the mode of each column specified in col for each group of a GroupBy object; sd calculates the standard deviation of each column specified in col for each group of a GroupBy object; ss calculates the sum of squares of each column specified in col for each group of a GroupBy object; sum calculates the sum of each column specified in col for each group of a GroupBy object; and var calculates the variance of each column specified in col for each group of a GroupBy object. If an aggregate is provided without a value (for example, as max in sum(col="X1", na="all").mean(col="X5", na="all").max()),
then it is assumed that the aggregation should apply to all columns except the GroupBy columns. However, operations will not be performed on String columns. They will be skipped. Note again that nrow is required and cannot be empty.

**Value**

Returns a new H2OFrame object with columns equivalent to the number of groups created

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
df <- h2o.importFile(paste("https://s3.amazonaws.com/h2o-public-test-data", 
"/smalldata/prostate/prostate.csv", 
sep=""))
h2o.group_by(data = df, by = "RACE", nrow("VOL"))
## End(Not run)
```

---

**h2o.gsub**

*String Global Substitute*

**Description**

Creates a copy of the target column in which each string has all occurrence of the regex pattern replaced with the replacement substring.

**Usage**

```r
h2o.gsub(pattern, replacement, x, ignore.case = FALSE)
```

**Arguments**

- `pattern`: The pattern to replace.
- `replacement`: The replacement pattern.
- `x`: The column on which to operate.
- `ignore.case`: Case sensitive or not

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
string_to_gsub <- as.h2o("r tutorial")
sub_string <- h2o.gsub("r ", "H2O ", string_to_gsub)
## End(Not run)
```
h2o.h Calulates Friedman and Popescu’s $H$ statistics, in order to test for the presence of an interaction between specified variables in h2o gbm and xgb models. $H$ varies from 0 to 1. It will have a value of 0 if the model exhibits no interaction between specified variables and a correspondingly larger value for a stronger interaction effect between them. NaN is returned if a computation is spoiled by weak main effects and rounding errors.

Description


Usage

h2o.h(model, frame, variables)

Arguments

- model: A trained gradient-boosting model.
- frame: A frame that current model has been fitted to.
- variables: Variables of the interest.

Examples

```r
## Not run:
library(h2o)
h2o.init()
prostate.hex <- h2o.importFile(
  destination_frame="prostate.hex"
)
prostate.hex$CAPSULE <- as.factor(prostate.hex$CAPSULE)
prostate.hex$RACE <- as.factor(prostate.hex$RACE)
prostate.h2o <- h2o.gbm(x = 3:9, y = "CAPSULE", training_frame = prostate.hex,
  distribution = "bernoulli", ntrees = 100, max_depth = 5, min_rows = 10, learn_rate = 0.1)
h_val <- h2o.h(prostate.h2o, prostate.hex, c("DPROS","DCAPS"))
## End(Not run)
```
h2o.head

Return the Head or Tail of an H2O Dataset.

Description

Returns the first or last rows of an H2OFrame object.

Usage

h2o.head(x, n = 6L, m = 200L, ...)  

## S3 method for class 'H2OFrame'
head(x, n = 6L, m = 200L, ...)  

h2o.tail(x, n = 6L, m = 200L, ...)  

## S3 method for class 'H2OFrame'
tail(x, n = 6L, m = 200L, ...)  

Arguments

x  
An H2OFrame object.

n  
(Optional) A single integer. If positive, number of rows in x to return. If negative, all but the n first/last number of rows in x.

m  
(Optional) A single integer. If positive, number of columns in x to return. If negative, all but the m first/last number of columns in x.

...  
Ignored.

Value

An H2OFrame containing the first or last n rows and m columns of an H2OFrame object.

Examples

## Not run:
library(h2o)
h2o.init(ip <- "localhost", port = 54321, startH2O = TRUE)
australia_path <- system.file("extdata", "australia.csv", package = "h2o")
australia <- h2o.uploadFile(path = australia_path)

# Return the first 10 rows and 6 columns
h2o.head(australia, n = 10L, m = 6L)

# Return the last 10 rows and 6 columns
h2o.tail(australia, n = 10L, m = 6L)

# For Jupyter notebook with an R kernel,
# view all rows of a data frame
options(repr.matrix.max.rows = 600, repr.matrix.max.cols = 200)

## End(Not run)
**h2o.HGLMMDetrics**

*Retrieve HGLM Model Metrics*

**Description**
Retrieve HGLM Model Metrics

**Usage**

```r
h2o.HGLMMDetrics(object)
```

**Arguments**

- `object` an H2OModel object or H2OModelMetrics.

---

**h2o.hist**

*Compute A Histogram*

**Description**
Compute a histogram over a numeric column. If breaks="FD", the MAD is used over the IQR in computing bin width. Note that we do not beautify the breakpoints as R does.

**Usage**

```r
h2o.hist(x, breaks = "Sturges", plot = TRUE)
```

**Arguments**

- `x` A single numeric column from an H2OFrame.
- `breaks` Can be one of the following: A string: "Sturges", "Rice", "sqrt", "Doane", "FD", "Scott" A single number for the number of breaks splitting the range of the vec into number of breaks bins of equal width A vector of numbers giving the split points, e.g., c(-50,213.2123,9324834)
- `plot` A logical value indicating whether or not a plot should be generated (default is TRUE).

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

iris <- h2o.importFile(f)
h2o.asnumeric(iris["petal_len"])
```
h2o.hit_ratio_table

Retrieve the Hit Ratios

Description

If "train", "valid", and "xval" parameters are FALSE (default), then the training Hit Ratios value is returned. If more than one parameter is set to TRUE, then a named list of Hit Ratio tables are returned, where the names are "train", "valid" or "xval".

Usage

h2o.hit_ratio_table(object, train = FALSE, valid = FALSE, xval = FALSE)

Arguments

object  An H2OModel object.
train   Retrieve the training Hit Ratio
valid   Retrieve the validation Hit Ratio
xval    Retrieve the cross-validation Hit Ratio

Examples

## Not run:
library(h2o)
h2o.init()

iris <- h2o.importFile(f)
iris_split <- h2o.splitFrame(data = iris, ratios = 0.8, seed = 1234)
train <- iris_split[[1]]
valid <- iris_split[[2]]

iris_xgb <- h2o.xgboost(x = 1:4, y = 5, training_frame = train, validation_frame = valid)
hrt_iris <- h2o.hit_ratio_table(iris_xgb, valid = TRUE)
hrt_iris

## End(Not run)
h2o.hour

Convert Milliseconds to Hour of Day in H2O Datasets

Description
Converts the entries of an H2OFrame object from milliseconds to hours of the day (on a 0 to 23 scale).

Usage
h2o.hour(x)

hour(x)

## S3 method for class 'H2OFrame'
hour(x)

Arguments
x An H2OFrame object.

Value
An H2OFrame object containing the entries of x converted to hours of the day.

See Also
h2o.day

h2o.ice_plot

Plot Individual Conditional Expectation (ICE) for each decile

Description
Individual Conditional Expectation (ICE) plot gives a graphical depiction of the marginal effect of a variable on the response. ICE plots are similar to partial dependence plots (PDP); PDP shows the average effect of a feature while ICE plot shows the effect for a single instance. This function will plot the effect for each decile. In contrast to the PDP, ICE plots can provide more insight, especially when there is stronger feature interaction.

Usage
h2o.ice_plot(model, newdata, column, target = NULL, max_levels = 30)
**h2o.ifelse**

**H2O Apply Conditional Statement**

**Description**

Applies conditional statements to numeric vectors in H2O parsed data objects when the data are numeric.

**Arguments**

- **model** (An H2OModel)
- **newdata** (An H2OFrame)
- **column** (A feature column name to inspect)
- **target** (If multinomial, plot PDP just for target category. Character string)
- **max_levels** (An integer specifying the maximum number of factor levels to show. Defaults to 30)

**Value**

A ggplot2 object

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

# Import the wine dataset into H2O:
df <- h2o.importFile(f)

# Set the response
response <- "quality"

# Split the dataset into a train and test set:
splits <- h2o.splitFrame(df, ratios = 0.8, seed = 1)
train <- splits[[1]]
test <- splits[[2]]

# Build and train the model:
gbm <- h2o.gbm(y = response, training_frame = train)

data <- h2o.gbm(y = response, training_frame = train)

# Create the individual conditional expectations plot
ice <- h2o.ice_plot(gbm, test, column = "alcohol")
print(ice)

## End(Not run)
```
h2o.ifelse(test, yes, no)

Arguments

test
A logical description of the condition to be met (>, <, =, etc...)
yes
The value to return if the condition is TRUE.
no
The value to return if the condition is FALSE.

Details

Both numeric and categorical values can be tested. However when returning a yes and no condition both conditions must be either both categorical or numeric.

Value

Returns a vector of new values matching the conditions stated in the ifelse call.

Examples

```r
## Not run:
library(h2o)
h2o.init()
australia_path <- system.file("extdata", "australia.csv", package = "h2o")
australia <- h2o.importFile(path = australia_path)
australia[, 9] <- ifelse(australia[, 3] < 279.9, 1, 0)
summary(australia)
## End(Not run)
```

---

h2o.importFile

Import Files into H2O

Description

Imports files into an H2O cluster. The default behavior is to pass-through to the parse phase automatically.

Usage

```r
h2o.importFile(
  path,
  destination_frame = "",
  parse = TRUE,
  header = NA,
```
h2o.importFile

sep = "",
col.names = NULL,
col.types = NULL,
na.strings = NULL,
decrypt_tool = NULL,
skipped_columns = NULL,
custom_non_data_line_markers = NULL,
partition_by = NULL,
quotechar = NULL,
escapechar = ""
)

h2o.importFolder(
    path,
    pattern = "", 
    destination_frame = "", 
    parse = TRUE, 
    header = NA, 
    sep = "", 
    col.names = NULL, 
    col.types = NULL, 
    na.strings = NULL, 
    decrypt_tool = NULL, 
    skipped_columns = NULL, 
    custom_non_data_line_markers = NULL, 
    partition_by = NULL, 
    quotechar = NULL, 
    escapechar = "\"
)

h2o.importHDFS(
    path,
    pattern = "", 
    destination_frame = "", 
    parse = TRUE, 
    header = NA, 
    sep = "", 
    col.names = NULL, 
    na.strings = NULL
)

h2o.uploadFile(
    path,
    destination_frame = "", 
    parse = TRUE, 
    header = NA, 
    sep = "", 
    col.names = NULL,
Arguments

path
The complete URL or normalized file path of the file to be imported. Each row of data appears as one line of the file.

destination_frame
(Optional) The unique hex key assigned to the imported file. If none is given, a key will automatically be generated based on the URL path.

parse
(Optional) A logical value indicating whether the file should be parsed after import, for details see h2o.parseRaw.

header
(Optional) A logical value indicating whether the first line of the file contains column headers. If left empty, the parser will try to automatically detect this.

sep
(Optional) The field separator character. Values on each line of the file are separated by this character. If sep = "", the parser will automatically detect the separator.

col.names
(Optional) An H2OFrame object containing a single delimited line with the column names for the file.

col.types
(Optional) A vector to specify whether columns should be forced to a certain type upon import parsing.

na.strings
(Optional) H2O will interpret these strings as missing.

decrypt_tool
(Optional) Specify a Decryption Tool (key-reference acquired by calling h2o.decryptionSetup.

skipped_columns
a list of column indices to be skipped during parsing.

custom_non_data_line_markers
(Optional) If a line in imported file starts with any character in given string it will NOT be imported. Empty string means all lines are imported, NULL means that default behaviour for given format will be used

partition_by
names of the columns the persisted dataset has been partitioned by.

quotechar
A hint for the parser which character to expect as quoting character. None (default) means autodetection.

escapechar
(Optional) One ASCII character used to escape other characters.

pattern
(Optional) Character string containing a regular expression to match file(s) in the folder.

progressBar
(Optional) When FALSE, tell H2O parse call to block synchronously instead of polling. This can be faster for small datasets but loses the progress bar.

parse_type
(Optional) Specify which parser type H2O will use. Valid types are "ARFF", "XLS", "CSV", "SVMLight"
**Details**

`h2o.importFile` is a parallelized reader and pulls information from the server from a location specified by the client. The path is a server-side path. This is a fast, scalable, highly optimized way to read data. H2O pulls the data from a data store and initiates the data transfer as a read operation.

Unlike the import function, which is a parallelized reader, `h2o.uploadFile` is a push from the client to the server. The specified path must be a client-side path. This is not scalable and is only intended for smaller data sizes. The client pushes the data from a local filesystem (for example, on your machine where R is running) to H2O. For big-data operations, you don’t want the data stored on or flowing through the client.

`h2o.importFolder` imports an entire directory of files. If the given path is relative, then it will be relative to the start location of the H2O instance. The default behavior is to pass-through to the parse phase automatically.

`h2o.importHDFS` is deprecated. Instead, use `h2o.importFile`.

**See Also**

`h2o.import_sql_select, h2o.import_sql_table, h2o.parseRaw`

**Examples**

```r
## Not run:
h2o.init(ip = "localhost", port = 54321, startH2O = TRUE)
prostate_path = system.file("extdata", "prostate.csv", package = "h2o")
prostate = h2o.importFile(path = prostate_path)
class(prostate)
summary(prostate)

#Import files with a certain regex pattern by utilizing h2o.importFolder()
#In this example we import all .csv files in the directory prostate_folder
prostate_path = system.file("extdata", "prostate_folder", package = "h2o")
prostate_pattern = h2o.importFolder(path = prostate_path, pattern = ".*.csv")
class(prostate_pattern)
summary(prostate_pattern)
## End(Not run)
```

---

**Description**

Import Hive table to H2OFrame in memory. Make sure to start H2O with Hive on classpath. Uses hive-site.xml on classpath to connect to Hive. When database is specified as jdbc URL uses Hive JDBC driver to obtain table metadata. then uses direct HDFS access to import data.
import_hive_table

Usage

h2o.import_hive_table(
    database,
    table,
    partitions = NULL,
    allow_multi_format = FALSE
)

Arguments

database         Name of Hive database (default database will be used by default), can be also a JDBC URL

name of Hive table to import

partitions       a list of lists of strings - partition key column values of partitions you want to import.

allow_multi_format enable import of partitioned tables with different storage formats used. WARNING: this may fail on out-of-memory for tables with a large number of small partitions.

Details

For example, my_citibike_data = h2o.import_hive_table("default", "citibike20k", partitions = list(c("2017", "01"), c("2017", "02"))) my_citibike_data = h2o.import_hive_table("jdbc:hive2://hive-server:10000/default", "citibike20k", allow_multi_format = TRUE)

-------------------

import_mojo

Imports a MOJO under given path, creating a Generic model with it.

Description

Usage example: mojo_model <- h2o.import_mojo(model_file_path = "/path/to/mojo.zip") predictions <- h2o.predict(mojo_model, dataset)

Usage

h2o.import_mojo(mojo_file_path, model_id = NULL)

Arguments

mojo_file_path Filesystem path to the model imported

model_id         Model ID, default is NULL

Value

Returns H2O Generic Model embedding given MOJO model
Examples

```r
## Not run:

# Import default Iris dataset as H2O frame
data <- as.h2o(iris)

# Train a very simple GBM model
original_model <- h2o.gbm(x = features, y = "Species", training_frame = data)

# Download the trained GBM model as MOJO (temporary directory used in this example)
mojo_original_path <- h2o.save_mojo(original_model, path = tempdir())

# Import the MOJO and obtain a Generic model
mojo_model <- h2o.import_mojo(mojo_original_path)

# Perform scoring with the generic model
predictions <- h2o.predict(mojo_model, data)

## End(Not run)
```

---

**h2o.import_sql_select**  
Import SQL table that is result of SELECT SQL query into H2O

---

**Description**

Creates a temporary SQL table from the specified sql_query. Runs multiple SELECT SQL queries on the temporary table concurrently for parallel ingestion, then drops the table. Be sure to start the h2o.jar in the terminal with your downloaded JDBC driver in the classpath: `java -cp <path_to_h2o_jar>:<path_to_jdbc_driver_jar> water.H2OApp` Also see h2o.import_sql_table. Currently supported SQL databases are MySQL, PostgreSQL, MariaDB, Hive, Oracle and Microsoft SQL Server.

**Usage**

```r
h2o.import_sql_select(
  connection_url,
  select_query,
  username,
  password,
  use_temp_table = NULL,
  temp_table_name = NULL,
  optimize = NULL,
  fetch_mode = NULL
)
```
Arguments

- **connection_url**: URL of the SQL database connection as specified by the Java Database Connectivity (JDBC) Driver. For example, "jdbc:mysql://localhost:3306/menagerie?useSSL=false"
- **select_query**: SQL query starting with 'SELECT' that returns rows from one or more database tables.
- **username**: Username for SQL server
- **password**: Password for SQL server
- **use_temp_table**: Whether a temporary table should be created from select_query
- **temp_table_name**: Name of temporary table to be created from select_query
- **optimize**: (Optional) Optimize import of SQL table for faster imports. Experimental. Default is true.
- **fetch_mode**: (Optional) Set to DISTRIBUTED to enable distributed imports. Set to SINGLE to force a sequential read from the database. Can be used for databases that do not support OFFSET-like clauses in SQL statements.

Details

For example, `my_sql_conn_url <- "jdbc:mysql://172.16.2.178:3306/ingestSQL?useSSL=false"`
`select_query <- "SELECT bikeid from citibike20k"`
`username <- "root"`
`password <- "abc123"`
`my_citibike_data <- h2o.import_sql_select(my_sql_conn_url, select_query, username, password)`

**Description**

Imports SQL table into an H2O cluster. Assumes that the SQL table is not being updated and is stable. Runs multiple SELECT SQL queries concurrently for parallel ingestion. Be sure to start the h2o.jar in the terminal with your downloaded JDBC driver in the classpath: `java -cp <path_to_h2o_jar>:<path_to_jdbc_driver_jar> water.H2OApp` Also see `h2o.import_sql_select`. Currently supported SQL databases are MySQL, PostgreSQL, MariaDB, Hive, Oracle and Microsoft SQL Server.

**Usage**

```r
h2o.import_sql_table(
  connection_url,  
  table,           
  username,       
  password,       
  columns = NULL, 
  optimize = NULL,  
  fetch_mode = NULL
)```
Arguments

classification_url  URL of the SQL database connection as specified by the Java Database Connectivity (JDBC) Driver. For example, "jdbc:mysql://localhost:3306/menagerie?&useSSL=false"

table  Name of SQL table
username  Username for SQL server
password  Password for SQL server

columns  (Optional) Character vector of column names to import from SQL table. Default is to import all columns.
optimize  (Optional) Optimize import of SQL table for faster imports. Default is true. Ignored - use fetch_mode instead.
fetch_mode  (Optional) Set to DISTRIBUTED to enable distributed import. Set to SINGLE to force a sequential read from the database. Can be used for databases that do not support OFFSET-like clauses in SQL statements.

Details

For example,

```
my_sql_conn_url <- "jdbc:mysql://172.16.2.178:3306/ingestSQL?&useSSL=false"
table <- "citibike20k" username <- "root" password <- "abc123" my_citibike_data <- h2o.import_sql_table(my_sql_conn_url, table, username, password)
```

Usage

```
h2o.impute(  
data,  
  column = 0,  
  method = c("mean", "median", "mode"),  
  combine_method = c("interpolate", "average", "lo", "hi"),  
  by = NULL,  
  groupByframe = NULL,  
  values = NULL  
)
```
Arguments

**data**
The dataset containing the column to impute.

**column**
A specific column to impute, default of 0 means impute the whole frame.

**method**
"mean" replaces NAs with the column mean; "median" replaces NAs with the column median; "mode" replaces with the most common factor (for factor columns only);

**combine_method**
If method is "median", then choose how to combine quantiles on even sample sizes. This parameter is ignored in all other cases.

**by**
group by columns

**groupByFrame**
Impute the column col with this pre-computed grouped frame.

**values**
A vector of impute values (one per column). NaN indicates to skip the column

Details

The default method is selected based on the type of the column to impute. If the column is numeric then "mean" is selected; if it is categorical, then "mode" is selected. Other column types (e.g. String, Time, UUID) are not supported.

Value

an H2OFrame with imputed values

Examples

```r
## Not run:
h2o.init()
iris_hf <- as.h2o(iris)
iris_hf[sample(nrow(iris_hf), 40), 5] <- NA # randomly replace 50 values with NA
# impute with a group by
iris_hf <- h2o.impute(iris_hf, "Species", "mode", by = c("Sepal.Length", "Sepal.Width"))
```

Description

The infogram is a graphical information-theoretic interpretability tool which allows the user to quickly spot the core, decision-making variables that uniquely and safely drive the response, in supervised classification problems. The infogram can significantly cut down the number of predictors needed to build a model by identifying only the most valuable, admissible features. When protected variables such as race or gender are present in the data, the admissibility of a variable is determined by a safety and relevancy index, and thus serves as a diagnostic tool for fairness. The safety of each feature can be quantified and variables that are unsafe will be considered inadmissible. Models built using only admissible features will naturally be more interpretable, given the reduced feature set. Admissible models are also less susceptible to overfitting and train faster, while providing similar accuracy as models built using all available features.
Usage

```r
h2o.infogram(
  x,
  y,
  training_frame,
  model_id = NULL,
  validation_frame = NULL,
  seed = -1,
  keep_cross_validation_models = TRUE,
  keep_cross_validation_predictions = FALSE,
  keep_cross_validation_fold_assignment = FALSE,
  nfolds = 0,
  fold_assignment = c("AUTO", "Random", "Modulo", "Stratified"),
  fold_column = NULL,
  ignore_const_cols = TRUE,
  score_each_iteration = FALSE,
  offset_column = NULL,
  weights_column = NULL,
  standardize = FALSE,
  distribution = c("AUTO", "bernoulli", "multinomial", "gaussian", "poisson", "gamma",
                   "tweedie", "laplace", "quantile", "huber"),
  plug_values = NULL,
  max_iterations = 0,
  stopping_rounds = 0,
  stopping_metric = c("AUTO", "deviance", "logloss", "MSE", "RMSE", "MAE", "RMSLE",
                      "AUC", "AUCPR", "lift_top_group", "misclassification", "mean_per_class_error",
                      "custom", "custom_increasing"),
  stopping_tolerance = 0.001,
  balance_classes = FALSE,
  class_sampling_factors = NULL,
  max_after_balance_size = 5,
  max_runtime_secs = 0,
  custom_metric_func = NULL,
  auc_type = c("AUTO", "NONE", "MACRO_OVR", "WEIGHTED_OVR", "MACRO_OVO",
              "WEIGHTED_OVO"),
  algorithm = c("AUTO", "deelearning", "drf", "gbm", "glm", "xgboost"),
  algorithm_params = NULL,
  protected_columns = NULL,
  total_information_threshold = -1,
  net_information_threshold = -1,
  relevance_index_threshold = -1,
  safety_index_threshold = -1,
  data_fraction = 1,
  top_n_features = 50
)
```
Arguments

x  (Optional) A vector containing the names or indices of the predictor variables to use in building the model. If x is missing, then all columns except y are used.

y  The name or column index of the response variable in the data. The response must be either a numeric or a categorical/factor variable. If the response is numeric, then a regression model will be trained, otherwise it will train a classification model.

training_frame  Id of the training data frame.

model_id  Destination id for this model; auto-generated if not specified.

validation_frame  Id of the validation data frame.

seed  Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).

keep_cross_validation_models  Logical. Whether to keep the cross-validation models. Defaults to TRUE.

keep_cross_validation_predictions  Logical. Whether to keep the predictions of the cross-validation models. Defaults to FALSE.

keep_cross_validation_fold_assignment  Logical. Whether to keep the cross-validation fold assignment. Defaults to FALSE.

nfolds  Number of folds for K-fold cross-validation (0 to disable or >= 2). Defaults to 0.

fold_assignment  Cross-validation fold assignment scheme, if fold_column is not specified. The 'Stratified' option will stratify the folds based on the response variable, for classification problems. Must be one of: "AUTO", "Random", "Modulo", "Stratified". Defaults to AUTO.

fold_column  Column with cross-validation fold index assignment per observation.

ignore_const_cols  Logical. Ignore constant columns. Defaults to TRUE.

score_each_iteration  Logical. Whether to score during each iteration of model training. Defaults to FALSE.

offset_column  Offset column. This will be added to the combination of columns before applying the link function.

weights_column  Column with observation weights. Giving some observation a weight of zero is equivalent to excluding it from the dataset; giving an observation a relative weight of 2 is equivalent to repeating that row twice. Negative weights are not allowed. Note: Weights are per-row observation weights and do not increase the size of the data frame. This is typically the number of times a row is repeated, but non-integer values are supported as well. During training, rows with higher weights matter more, due to the larger loss function pre-factor. If you set weight = 0 for a row, the returned prediction frame at that row is zero and this is incorrect. To get an accurate prediction, remove all rows with weight == 0.
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>standardize</td>
<td>Logical. Standardize numeric columns to have zero mean and unit variance. Defaults to FALSE.</td>
</tr>
<tr>
<td>distribution</td>
<td>Distribution function Must be one of: &quot;AUTO&quot;, &quot;bernoulli&quot;, &quot;multinomial&quot;, &quot;gaussian&quot;, &quot;poisson&quot;, &quot;gamma&quot;, &quot;tweedie&quot;, &quot;laplace&quot;, &quot;quantile&quot;, &quot;huber&quot;. Defaults to AUTO.</td>
</tr>
<tr>
<td>plug_values</td>
<td>Plug Values (a single row frame containing values that will be used to impute missing values of the training/validation frame, use with conjunction missing_values_handling = PlugValues).</td>
</tr>
<tr>
<td>max_iterations</td>
<td>Maximum number of iterations. Defaults to 0.</td>
</tr>
<tr>
<td>stopping_rounds</td>
<td>Early stopping based on convergence of stopping_metric. Stop if simple moving average of length k of the stopping_metric does not improve for k:=stopping_rounds scoring events (0 to disable) Defaults to 0.</td>
</tr>
<tr>
<td>stopping_metric</td>
<td>Metric to use for early stopping (AUTO: logloss for classification, deviance for regression and anomaly_score for Isolation Forest). Note that custom and custom_increasing can only be used in GBM and DRF with the Python client. Must be one of: &quot;AUTO&quot;, &quot;deviance&quot;, &quot;logloss&quot;, &quot;MSE&quot;, &quot;RMSE&quot;, &quot;MAE&quot;, &quot;RMSLE&quot;, &quot;AUC&quot;, &quot;AUCPR&quot;, &quot;lift_top_group&quot;, &quot;misclassification&quot;, &quot;mean_per_class_error&quot;, &quot;custom&quot;, &quot;custom_increasing&quot;. Defaults to AUTO.</td>
</tr>
<tr>
<td>stopping_tolerance</td>
<td>Relative tolerance for metric-based stopping criterion (stop if relative improvement is not at least this much) Defaults to 0.001.</td>
</tr>
<tr>
<td>balance_classes</td>
<td>Logical. Balance training data class counts via over/under-sampling (for imbalanced data). Defaults to FALSE.</td>
</tr>
<tr>
<td>class_sampling_factors</td>
<td>Desired over/under-sampling ratios per class (in lexicographic order). If not specified, sampling factors will be automatically computed to obtain class balance during training. Requires balance_classes.</td>
</tr>
<tr>
<td>max_after_balance_size</td>
<td>Maximum relative size of the training data after balancing class counts (can be less than 1.0). Requires balance_classes. Defaults to 5.0.</td>
</tr>
<tr>
<td>max_runtime_secs</td>
<td>Maximum allowed runtime in seconds for model training. Use 0 to disable. Defaults to 0.</td>
</tr>
<tr>
<td>custom_metric_func</td>
<td>Reference to custom evaluation function, format: ‘language:keyName=funcName’</td>
</tr>
<tr>
<td>auc_type</td>
<td>Set default multinomial AUC type. Must be one of: &quot;AUTO&quot;, &quot;NONE&quot;, &quot;MACRO_OVR&quot;, &quot;WEIGHTED_OVR&quot;, &quot;MACRO_OVO&quot;, &quot;WEIGHTED_OVO&quot;. Defaults to AUTO.</td>
</tr>
<tr>
<td>algorithm</td>
<td>Type of machine learning algorithm used to build the infogram. Options include 'AUTO' (gbm), 'deeplearning' (Deep Learning with default parameters), 'drf' (Random Forest with default parameters), 'gbm' (GBM with default parameters), 'glm' (GLM with default parameters), or 'xgboost' (if available, XGBoost with default parameters). Must be one of: &quot;AUTO&quot;, &quot;deeplearning&quot;, &quot;drf&quot;, &quot;gbm&quot;, &quot;glm&quot;, &quot;xgboost&quot;. Defaults to AUTO.</td>
</tr>
</tbody>
</table>
algorithm_params
Customized parameters for the machine learning algorithm specified in the algorithm parameter.

protected_columns
Columns that contain features that are sensitive and need to be protected (legally, or otherwise), if applicable. These features (e.g. race, gender, etc) should not drive the prediction of the response.

total_information_threshold
A number between 0 and 1 representing a threshold for total information, defaulting to 0.1. For a specific feature, if the total information is higher than this threshold, and the corresponding net information is also higher than the threshold “net_information_threshold”, that feature will be considered admissible. The total information is the x-axis of the Core Infogram. Default is -1 which gets set to 0.1. Defaults to -1.

net_information_threshold
A number between 0 and 1 representing a threshold for net information, defaulting to 0.1. For a specific feature, if the net information is higher than this threshold, and the corresponding total information is also higher than the total_information_threshold, that feature will be considered admissible. The net information is the y-axis of the Core Infogram. Default is -1 which gets set to 0.1. Defaults to -1.

relevance_index_threshold
A number between 0 and 1 representing a threshold for the relevance index, defaulting to 0.1. This is only used when “protected_columns” is set by the user. For a specific feature, if the relevance index value is higher than this threshold, and the corresponding safety index is also higher than the safety_index_threshold, that feature will be considered admissible. The relevance index is the x-axis of the Fair Infogram. Default is -1 which gets set to 0.1. Defaults to -1.

safety_index_threshold
A number between 0 and 1 representing a threshold for the safety index, defaulting to 0.1. This is only used when protected_columns is set by the user. For a specific feature, if the safety index value is higher than this threshold, and the corresponding relevance index is also higher than the relevance_index_threshold, that feature will be considered admissible. The safety index is the y-axis of the Fair Infogram. Default is -1 which gets set to 0.1. Defaults to -1.

data_fraction
The fraction of training frame to use to build the infogram model. Defaults to 1.0, and any value greater than 0 and less than or equal to 1.0 is acceptable. Defaults to 1.

top_n_features
An integer specifying the number of columns to evaluate in the infogram. The columns are ranked by variable importance, and the top N are evaluated. Defaults to 50. Defaults to 50.

Details
The infogram allows the user to quickly spot the admissible decision-making variables that are driving the response. There are two types of infogram plots: Core and Fair Infogram.

The Core Infogram plots all the variables as points on two-dimensional grid of total vs net information. The x-axis is total information, a measure of how much the variable drives the response (the
more predictive, the higher the total information). The y-axis is net information, a measure of how unique the variable is. The top right quadrant of the infogram plot is the admissible section; the variables located in this quadrant are the admissible features. In the Core Infogram, the admissible features are the strongest, unique drivers of the response.

If sensitive or protected variables are present in data, the user can specify which attributes should be protected while training using the `protected_columns` argument. All non-protected predictor variables will be checked to make sure that there’s no information pathway to the response through a protected feature, and deemed inadmissible if they possess little or no informational value beyond their use as a dummy for protected attributes. The Fair Infogram plots all the features as points on two-dimensional grid of relevance vs safety. The x-axis is relevance index, a measure of how much the variable drives the response (the more predictive, the higher the relevance). The y-axis is safety index, a measure of how much extra information the variable has that is not acquired through the protected variables. In the Fair Infogram, the admissible features are the strongest, safest drivers of the response.

Examples

```r
## Not run:
h2o.init()

# Convert iris dataset to an H2OFrame
df <- as.h2o(iris)

# Infogram
ig <- h2o.infogram(y = "Species", training_frame = df)
plot(ig)

## End(Not run)
```

### h2o.init

##### Initialize and Connect to H2O

Attempts to start and/or connect to and H2O instance.

#### Usage

```r
h2o.init(
  ip = "localhost",
  port = 54321,
  name = NA_character_,
  startH2O = TRUE,
  forceDL = FALSE,
  enable_assertions = TRUE,
  license = NULL,
  nthreads = -1,
)```
max_mem_size = NULL,
min_mem_size = NULL,
ice_root = tempdir(),
log_dir = NA_character_,
log_level = NA_character_,
strict_version_check = TRUE,
proxy = NA_character_,
https = FALSE,
cacert = NA_character_,
insecure = FALSE,
username = NA_character_,
password = NA_character_,
use_spnego = FALSE,
cookies = NA_character_,
context_path = NA_character_,
ignore_config = FALSE,
extra_classpath = NULL,
jvm_custom_args = NULL,
bind_to_localhost = TRUE
)

Arguments

ip Object of class character representing the IP address of the server where H2O is running.
port Object of class numeric representing the port number of the H2O server.
nname (Optional) A character string representing the H2O cluster name.
startH2O (Optional) A logical value indicating whether to try to start H2O from R if no connection with H2O is detected. This is only possible if ip = "localhost" or ip = "127.0.0.1". If an existing connection is detected, R does not start H2O.
forceDL (Optional) A logical value indicating whether to force download of the H2O executable. Defaults to FALSE, so the executable will only be downloaded if it does not already exist in the h2o R library resources directory h2o/java/h2o.jar. This value is only used when R starts H2O.
enable_assertions (Optional) A logical value indicating whether H2O should be launched with assertions enabled. Used mainly for error checking and debugging purposes. This value is only used when R starts H2O.
license (Optional) A character string value specifying the full path of the license file. This value is only used when R starts H2O.
nthreads (Optional) Number of threads in the thread pool. This relates very closely to the number of CPUs used. -1 means use all CPUs on the host (Default). A positive integer specifies the number of CPUs directly. This value is only used when R starts H2O.
max_mem_size (Optional) A character string specifying the maximum size, in bytes, of the memory allocation pool to H2O. This value must a multiple of 1024 greater
than 2MB. Append the letter m or M to indicate megabytes, or g or G to indicate gigabytes. This value is only used when R starts H2O. If max_mem_size is not defined, then the amount of memory that H2O allocates will be determined by the default memory of Java Virtual Machine. This amount is dependent on the Java version, but it will generally be 25 percent of the machine’s physical memory.

**min_mem_size**  (Optional) A character string specifying the minimum size, in bytes, of the memory allocation pool to H2O. This value must a multiple of 1024 greater than 2MB. Append the letter m or M to indicate megabytes, or g or G to indicate gigabytes. This value is only used when R starts H2O.

**ice_root**  (Optional) A directory to handle object spillage. The default varies by OS.

**log_dir**  (Optional) A directory where H2O server logs are stored. The default varies by OS.

**log_level**  (Optional) The level of logging of H2O server. The default is INFO.

**strict_version_check**  (Optional) Setting this to FALSE is unsupported and should only be done when advised by technical support.

**proxy**  (Optional) A character string specifying the proxy path.

**https**  (Optional) Set this to TRUE to use https instead of http.

**cacert**  (Optional) Path to a CA bundle file with root and intermediate certificates of trusted CAs.

**insecure**  (Optional) Set this to TRUE to disable SSL certificate checking.

**username**  (Optional) Username to login with.

**password**  (Optional) Password to login with.

**use_spnego**  (Optional) Set this to TRUE to enable SPNEGO authentication.

**cookies**  (Optional) Vector(or list) of cookies to add to request.

**context_path**  (Optional) The last part of connection URL: http://<ip>:<port>/<context_path>

**ignore_config**  (Optional) A logical value indicating whether a search for a .h2oconfig file should be conducted or not. Default value is FALSE.

**extra_classpath**  (Optional) A vector of paths to libraries to be added to the Java classpath when H2O is started from R.

**jvm_custom_args**  (Optional) A character list of custom arguments for the JVM where new H2O instance is going to run, if started. Ignored when connecting to an existing instance.

**bind_to_localhost**  (Optional) A logical flag indicating whether access to the H2O instance should be restricted to the local machine (default) or if it can be reached from other computers on the network. Only applicable when H2O is started from R.
Details

By default, this method first checks if an H2O instance is connectible. If it cannot connect and start = TRUE with ip = "localhost", it will attempt to start and instance of H2O at localhost:54321. If an open ip and port of your choice are passed in, then this method will attempt to start an H2O instance at that specified ip port.

When initializing H2O locally, this method searches for h2o.jar in the R library resources (system.file("java","h2o.jar"), and if the file does not exist, it will automatically attempt to download the correct version from Amazon S3. The user must have Internet access for this process to be successful.

Once connected, the method checks to see if the local H2O R package version matches the version of H2O running on the server. If there is a mismatch and the user indicates she wishes to upgrade, it will remove the local H2O R package and download/install the H2O R package from the server.

Value

this method will load it and return a H2OConnection object containing the IP address and port number of the H2O server.

Note

Users may wish to manually upgrade their package (rather than waiting until being prompted), which requires that they fully uninstall and reinstall the H2O package, and the H2O client package. You must unload packages running in the environment before upgrading. It’s recommended that users restart R or R studio after upgrading

See Also

H2O R package documentation for more details. h2o.shutdown for shutting down from R.

Examples

```r
## Not run:
# Try to connect to a local H2O instance that is already running.
# If not found, start a local H2O instance from R with the default settings.
h2o.init()

# Try to connect to a local H2O instance.
# If not found, raise an error.
h2o.init(startH2O = FALSE)

# Try to connect to a local H2O instance that is already running.
# If not found, start a local H2O instance from R with 5 gigabytes of memory.
h2o.init(max_mem_size = "5g")

# Try to connect to a local H2O instance that is already running.
# If not found, start a local H2O instance from R that uses 5 gigabytes of memory.
h2o.init(max_mem_size = "5g")

## End(Not run)
```
h2o.insertMissingValues

*Insert Missing Values into an H2OFrame*

**Description**

Randomly replaces a user-specified fraction of entries in an H2O dataset with missing values.

**Usage**

```r
h2o.insertMissingValues(data, fraction = 0.1, seed = -1)
```

**Arguments**

- `data`:
  
  An H2OFrame object representing the dataset.

- `fraction`:
  
  A number between 0 and 1 indicating the fraction of entries to replace with missing.

- `seed`:
  
  A random number used to select which entries to replace with missing values. Default of seed = -1 will automatically generate a seed in H2O.

**Value**

Returns an H2OFrame object.

**WARNING**

This will modify the original dataset. Unless this is intended, this function should only be called on a subset of the original.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

iris_hf <- as.h2o(iris)
summary(iris_hf)

iris_miss <- h2o.insertMissingValues(iris_hf, fraction = 0.25)
head(iris_miss)
summary(iris_miss)

## End(Not run)
```
Description

Creates a data frame in H2O with n-th order interaction features between categorical columns, as specified by the user.

Usage

h2o.interaction(
  data,
  destination_frame,
  factors,
  pairwise,
  max_factors,
  min_occurrence
)

Arguments

data An H2OFrame object containing the categorical columns.
destination_frame A string indicating the destination key. If empty, this will be auto-generated by H2O.
factors Factor columns (either indices or column names).
pairwise Whether to create pairwise interactions between factors (otherwise create one higher-order interaction). Only applicable if there are 3 or more factors.
max_factors Max. number of factor levels in pair-wise interaction terms (if enforced, one extra catch-all factor will be made)
min_occurrence Min. occurrence threshold for factor levels in pair-wise interaction terms

Value

Returns an H2OFrame object.

Examples

## Not run:
library(h2o)
h2o.init()

# Create some random data
my_frame <- h2o.createFrame(rows = 20, cols = 5,
    seed = -12301283, randomize = TRUE, value = 0,
    categorical_fraction = 0.8, factors = 10, real_range = 1,
    integer_fraction = 0.2, integer_range = 10,
binary_fraction = 0, binary_ones_fraction = 0.5, 
missing_fraction = 0.2, 
response_factors = 1)

# Turn integer column into a categorical
my_frame[,5] <- as.factor(my_frame[,5])
head(my_frame, 20)

# Create pairwise interactions
pairwise <- h2o.interaction(my_frame, 
    factors = list(c(1, 2), c("C2", "C3", "C4")),
    pairwise = TRUE, max_factors = 10, min_occurrence = 1)
head(pairwise, 20)
h2o.levels(pairwise, 2)

# Create 5-th order interaction
highorder <- h2o.interaction(my_frame, factors = c(1, 2, 3, 4, 5),
    pairwise = FALSE, max_factors = 10000, min_occurrence = 1)
head(highorder, 20)

# Limit the number of factors of the "categoricalized" integer column
# to at most 3 factors, and only if they occur at least twice
head(my_frame[,5], 20)
trim_integer_levels <- h2o.interaction(my_frame, factors = "C5", pairwise = FALSE, max_factors = 3,
    min_occurrence = 2)
head(trim_integer_levels, 20)

# Put all together
my_frame <- h2o.cbind(my_frame, pairwise, higherorder, trim_integer_levels)
my_frame
head(my_frame, 20)
summary(my_frame)

## End(Not run)

---

### h2o.isax

**iSAX**

**Description**

Compute the iSAX index for a DataFrame which is assumed to be numeric time series data

**Usage**

```
h2o.isax(x, num_words, max_cardinality, optimize_card = FALSE)
```

**Arguments**

- **x** an H2OFrame
- **num_words** Number of iSAX words for the timeseries. ie granularity along the time series
h2o.ischaracter

**Description**
Check if character

**Usage**
h2o.ischaracter(x)

**Arguments**
x An H2OFrame object.

**See Also**
character for the base R implementation, is.character().
Examples

```r
## Not run:
library(h2o)
h2o.init()

iris <- h2o.importFile(f)
iris_char <- h2o.ascharacter(iris[["class"]])
h2o.ischaracter(iris_char)

## End(Not run)
```

h2o.isfactor

Check if factor

Description

Check if factor

Usage

h2o.isfactor(x)

Arguments

x

An H2OFrame object.

See Also

factor for the base R implementation, is.factor().

Examples

```r
## Not run:
library(h2o)
h2o.init()

cars <- h2o.importFile(f)
cars["economy_20mpg"] <- as.factor(cars["economy_20mpg"])
h2o.isfactor(cars["economy_20mpg"])

## End(Not run)
```
**h2o.isnumeric**  
*Check if numeric*

**Description**  
Check if numeric

**Usage**  
h2o.isnumeric(x)

**Arguments**  
x  
An H2OFrame object.

**See Also**  
numeric for the base R implementation, is.numeric().

**Examples**  
## Not run:  
library(h2o)  
h2o.init()

iris <- h2o.importFile(f)  
h2o.isnumeric(iris["sepal_len"])

## End(Not run)

---

**h2o.isolationForest**  
*Trains an Isolation Forest model*

**Description**  
Trains an Isolation Forest model

**Usage**  
h2o.isolationForest(  
training_frame,  
x,  
model_id = NULL,  
score_each_iteration = FALSE,  
score_tree_interval = 0,  
ignore_const_cols = TRUE,
ntrees = 50,
max_depth = 8,
min_rows = 1,
max_runtime_secs = 0,
seed = -1,
build_tree_one_node = FALSE,
mtries = -1,
sample_size = 256,
sample_rate = -1,
col_sample_rate_change_per_level = 1,
col_sample_rate_per_tree = 1,
categorical_encoding = c("AUTO", "Enum", "OneHotInternal", "OneHotExplicit",
  "Binary", "Eigen", "LabelEncoder", "SortByResponse", "EnumLimited"),
stopping_rounds = 0,
stopping_metric = c("AUTO", "anomaly_score"),
stopping_tolerance = 0.01,
export_checkpoints_dir = NULL,
contamination = -1,
validation_frame = NULL,
validation_response_column = NULL
}

Arguments

training_frame  Id of the training data frame.
x  A vector containing the character names of the predictors in the model.
model_id  Destination id for this model; auto-generated if not specified.
score_each_iteration  Logical. Whether to score during each iteration of model training. Defaults to FALSE.
score_tree_interval  Score the model after every so many trees. Disabled if set to 0. Defaults to 0.
ignore_const_cols  Logical. Ignore constant columns. Defaults to TRUE.
ntrees  Number of trees. Defaults to 50.
max_depth  Maximum tree depth (0 for unlimited). Defaults to 8.
min_rows  Fewest allowed (weighted) observations in a leaf. Defaults to 1.
max_runtime_secs  Maximum allowed runtime in seconds for model training. Use 0 to disable. Defaults to 0.
seed  Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).
build_tree_one_node  Logical. Run on one node only; no network overhead but fewer cpus used. Suitable for small datasets. Defaults to FALSE.
mtries  Number of variables randomly sampled as candidates at each split. If set to -1, defaults (number of predictors)/3. Defaults to -1.

sample_size  Number of randomly sampled observations used to train each Isolation Forest tree. Only one of parameters sample_size and sample_rate should be defined. If sample_rate is defined, sample_size will be ignored. Defaults to 256.

sample_rate  Rate of randomly sampled observations used to train each Isolation Forest tree. Needs to be in range from 0.0 to 1.0. If set to -1, sample_rate is disabled and sample_size will be used instead. Defaults to -1.

col_sample_rate_change_per_level  Relative change of the column sampling rate for every level (must be > 0.0 and <= 2.0) Defaults to 1.

col_sample_rate_per_tree  Column sample rate per tree (from 0.0 to 1.0) Defaults to 1.

categorical_encoding  Encoding scheme for categorical features Must be one of: "AUTO", "Enum", "OneHotInternal", "OneHotExplicit", "Binary", "Eigen", "LabelEncoder", "Sort-ByResponse", "EnumLimited". Defaults to AUTO.

stopping_rounds  Early stopping based on convergence of stopping_metric. Stop if simple moving average of length k of the stopping_metric does not improve for k:=stopping_rounds scoring events (0 to disable) Defaults to 0.

stopping_metric  Metric to use for early stopping (AUTO: logloss for classification, deviance for regression and anomaly_score for Isolation Forest). Note that custom and custom_increasing can only be used in GBM and DRF with the Python client. Must be one of: "AUTO", "anomaly_score". Defaults to AUTO.

stopping_tolerance  Relative tolerance for metric-based stopping criterion (stop if relative improvement is not at least this much) Defaults to 0.01.

export_checkpoints_dir  Automatically export generated models to this directory.

contamination  Contamination ratio - the proportion of anomalies in the input dataset. If undefined (-1) the predict function will not mark observations as anomalies and only anomaly score will be returned. Defaults to -1 (undefined). Defaults to -1.

validation_frame  Id of the validation data frame.

validation_response_column  (experimental) Name of the response column in the validation frame. Response column should be binary and indicate not anomaly/anomaly.

Examples

```r
# Not run:
library(h2o)
h2o.init()

# Import the cars dataset
```
cars <- h2o.importFile(f)

# Set the predictors
predictors <- c("displacement", "power", "weight", "acceleration", "year")

# Train the IF model
cars_if <- h2o.isolationForest(x = predictors, training_frame = cars,
   seed = 1234, stopping_metric = "anomaly_score",
   stopping_rounds = 3, stopping_tolerance = 0.1)

## End(Not run)

---

**h2o.is_client**

*Check Client Mode Connection*

**Description**

Check Client Mode Connection

**Usage**

h2o.is_client()

---

**h2o.keyof**

*Method on Keyed objects allowing to obtain their key.*

**Description**

Method on Keyed objects allowing to obtain their key.

**Usage**

h2o.keyof(object)

## S4 method for signature 'Keyed'
h2o.keyof(object)

## S4 method for signature 'H2OModel'
h2o.keyof(object)

## S4 method for signature 'H2OGrid'
h2o.keyof(object)

## S4 method for signature 'H2OFrame'
h2o.keyof(object)

## S4 method for signature 'H2OAutoML'
h2o.keyof(object)
h2o.kfold_column

Arguments

object A Keyed object

Value

the string key holding the persistent object.

Description

Create a k-fold vector useful for H2O algorithms that take a fold_assignments argument.

Usage

h2o.kfold_column(data, nfolds, seed = -1)

Arguments

data A dataframe against which to create the fold column.
nfolds The number of desired folds.
seed A random seed, -1 indicates that H2O will choose one.

Value

Returns an H2OFrame object with fold assignments.

Examples

## Not run:
library(h2o)
h2o.init()

iris <- h2o.importFile(f)
kfolds <- h2o.kfold_column(iris, nfolds = 5, seed = 1234)

## End(Not run)
**h2o.killMinus3**

*Dump the stack into the JVM’s stdout.*

**Description**

A poor man’s profiler, but effective.

**Usage**

```r
h2o.killMinus3()
```

**h2o.kmeans**

*Performs k-means clustering on an H2O dataset*

**Description**

Performs k-means clustering on an H2O dataset

**Usage**

```r
h2o.kmeans(
  training_frame,
  x,
  model_id = NULL,
  validation_frame = NULL,
  nfolds = 0,
  keep_cross_validation_models = TRUE,
  keep_cross_validation_predictions = FALSE,
  keep_cross_validation_fold_assignment = FALSE,
  fold_assignment = c("AUTO", "Random", "Modulo", "Stratified"),
  fold_column = NULL,
  ignore_const_cols = TRUE,
  score_each_iteration = FALSE,
  k = 1,
  estimate_k = FALSE,
  user_points = NULL,
  max_iterations = 10,
  standardize = TRUE,
  seed = -1,
  init = c("Random", "PlusPlus", "Furthest", "User"),
  max_runtime_secs = 0,
  categorical_encoding = c("AUTO", "Enum", "OneHotInternal", "OneHotExplicit",
                          "Binary", "Eigen", "LabelEncoder", "SortByResponse", "EnumLimited"),
  export_checkpoints_dir = NULL,
  cluster_size_constraints = NULL,
)
```
Arguments

training_frame  Id of the training data frame.
x  A vector containing the character names of the predictors in the model.
model_id  Destination id for this model; auto-generated if not specified.
validation_frame  Id of the validation data frame.
nfolds  Number of folds for K-fold cross-validation (0 to disable or >= 2). Defaults to 0.
keep_cross_validation_models  Logical. Whether to keep the cross-validation models. Defaults to TRUE.
keep_cross_validation_predictions  Logical. Whether to keep the predictions of the cross-validation models. Defaults to FALSE.
keep_cross_validation_fold_assignment  Logical. Whether to keep the cross-validation fold assignment. Defaults to FALSE.
fold_assignment  Cross-validation fold assignment scheme, if fold_column is not specified. The 'Stratified' option will stratify the folds based on the response variable, for classification problems. Must be one of: "AUTO", "Random", "Modulo", "Stratified". Defaults to AUTO.
fold_column  Column with cross-validation fold index assignment per observation.
ignore_const_cols  Logical. Ignore constant columns. Defaults to TRUE.
score_each_iteration  Logical. Whether to score during each iteration of model training. Defaults to FALSE.
k  The max. number of clusters. If estimate_k is disabled, the model will find k centroids, otherwise it will find up to k centroids. Defaults to 1.
estimate_k  Logical. Whether to estimate the number of clusters (<=k) iteratively and deterministically. Defaults to FALSE.
user_points  This option allows you to specify a dataframe, where each row represents an initial cluster center. The user-specified points must have the same number of columns as the training observations. The number of rows must equal the number of clusters
max_iterations  Maximum training iterations (if estimate_k is enabled, then this is for each inner Lloyds iteration) Defaults to 10.
standardize  Logical. Standardize columns before computing distances. Defaults to TRUE.
seed  Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).
max_runtime_secs
Maximum allowed runtime in seconds for model training. Use 0 to disable.
Defaults to 0.
categorical_encoding
Encoding scheme for categorical features. Must be one of: "AUTO", "Enum",
"OneHotInternal", "OneHotExplicit", "Binary", "Eigen", "LabelEncoder", "SortByResponse",
"EnumLimited". Defaults to AUTO.
export_checkpoints_dir
Automatically export generated models to this directory.
cluster_size_constraints
An array specifying the minimum number of points that should be in each cluster.
The length of the constraints array has to be the same as the number of clusters.

Value
an object of class H2OClusteringModel.

See Also
h2o.cluster_sizes, h2o.totss, h2o.num_iterations, h2o.betweenss, h2o.tot_withinss,
h2o.withinss, h2o.centersSTD, h2o.centers

Examples
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
h2o.kmeans(training_frame = prostate, k = 10, x = c("AGE", "RACE", "VOL", "GLEASON"))
## End(Not run)

h2o.kolmogorov_smirnov
Kolmogorov-Smirnov metric for binomial models

Description
Retrieves a Kolmogorov-Smirnov metric for given binomial model. The number returned is in
range between 0 and 1. K-S metric represents the degree of separation between the positive (1) and
negative (0) cumulative distribution functions. Detailed metrics per each group are to be found in
the gains-lift table.
Usage

h2o.kolmogorov_smirnov(object)

## S4 method for signature 'H2OModelMetrics'
h2o.kolmogorov_smirnov(object)

## S4 method for signature 'H2OModel'
h2o.kolmogorov_smirnov(object)

Arguments

object Either an H2OModel object or an H2OModelMetrics object.

Details

The H2OModelMetrics version of this function will only take H2OBinomialMetrics objects.

Value

Kolmogorov-Smirnov metric, a number between 0 and 1.

See Also

h2o.gainsLift to see detailed K-S metrics per group

Examples

## Not run:
library(h2o)
h2o.init()
data <- h2o.importFile(
  path = "https://s3.amazonaws.com/h2o-public-test-data/smalldata/airlines/allyears2k_headers.zip")
model <- h2o.gbm(x = c("Origin", "Distance"), y = "IsDepDelayed",
  training_frame = data, ntree = 1)
h2o.kolmogorov_smirnov(model)

## End(Not run)

---

h2o.kurtosis

Kurtosis of a column

Description

Obtain the kurtosis of a column of a parsed H2O data object.
Usage

h2o.kurtosis(x, ..., na.rm = TRUE)

kurtosis.H2OFrame(x, ..., na.rm = TRUE)

Arguments

x
An H2OFrame object.

... Further arguments to be passed from or to other methods.

na.rm A logical value indicating whether NA or missing values should be stripped before the computation.

Value

Returns a list containing the kurtosis for each column (NaN for non-numeric columns).

Examples

## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
h2o.kurtosis(prostate$AGE)
## End(Not run)

h2o.learning_curve_plot

Learning Curve Plot

Description

Create learning curve plot for an H2O Model. Learning curves show error metric dependence on learning progress, e.g., RMSE vs number of trees trained so far in GBM. There can be up to 4 curves showing Training, Validation, Training on CV Models, and Cross-validation error.

Usage

h2o.learning_curve_plot(
  model,
  metric = c("AUTO", "auc", "aucpr", "mae", "rmse", "anomaly_score", "convergence",
             "custom", "custom_increasing", "deviance", "lift_top_group", "logloss",
             "misclassification", "negative_log_likelihood", "objective", "summetaieta02"),
  cv_ribbon = NULL,
  cv_lines = NULL
)
Arguments

- `model`: an H2O model
- `metric`: Metric to be used for the learning curve plot. These should mostly correspond with stopping metric.
- `cv_ribbon`: if True, plot the CV mean as a and CV standard deviation as a ribbon around the mean, if NULL, it will attempt to automatically determine if this is suitable visualisation
- `cv_lines`: if True, plot scoring history for individual CV models, if NULL, it will attempt to automatically determine if this is suitable visualisation

Value

A ggplot2 object

Examples

```r
## Not run:
library(h2o)
h2o.init()

# Import the wine dataset into H2O:
df <- h2o.importFile(f)

# Set the response
response <- "quality"

# Split the dataset into a train and test set:
splits <- h2o.splitFrame(df, ratios = 0.8, seed = 1)
train <- splits[[1]]
test <- splits[[2]]

# Build and train the model:
gbm <- h2o.gbm(y = response,
              training_frame = train)

# Create the learning curve plot
learning_curve <- h2o.learning_curve_plot(gbm)
print(learning_curve)

## End(Not run)
```

h2o.levels

Return the levels from the column requested column.

Description

Return the levels from the column requested column.
Usage

h2o.levels(x, i)

Arguments

x
An H2OFrame object.

i
Optional, the index of the column whose domain is to be returned.

See Also

levels for the base R method.

Examples

## Not run:
library(h2o)
h2o.init()

iris_hf <- as.h2o(iris)
h2o.levels(iris_hf, 5) # returns "setosa"  "versicolor"  "virginica"

## End(Not run)

---

h2o.listTimezones  List all of the Time Zones Acceptable by the H2O cluster.

Description

List all of the Time Zones Acceptable by the H2O cluster.

Usage

h2o.listTimezones()

---

h2o.list_all_extensions  List all H2O registered extensions

Description

List all H2O registered extensions

Usage

h2o.list_all_extensions()
**h2o.list_api_extensions**

*List registered API extensions*

**Description**

List registered API extensions

**Usage**

```r
h2o.list_api_extensions()
```

**h2o.list_core_extensions**

*List registered core extensions*

**Description**

List registered core extensions

**Usage**

```r
h2o.list_core_extensions()
```

**h2o.list_jobs**

*Return list of jobs performed by the H2O cluster*

**Description**

Return list of jobs performed by the H2O cluster

**Usage**

```r
h2o.list_jobs()
```
h2o.list_models  
Get an list of all model ids present in the cluster

Description

Get an list of all model ids present in the cluster

Usage

h2o.list_models()

Value

Returns a vector of model ids.

Examples

## Not run:
library(h2o)
h2o.init()

iris_hf <- as.h2o(iris)
model_id <- h2o.gbm(x = 1:4, y = 5, training_frame = iris_hf)@model_id
model_id_list <- h2o.list_models()

## End(Not run)

h2o.loadGrid  
Loads previously saved grid with all it's models from the same folder

Description

Returns a reference to the loaded Grid.

Usage

h2o.loadGrid(grid_path, load_params_references = FALSE)

Arguments

grid_path  
A character string containing the path to the file with the grid saved.

load_params_references  
A logical which if true will attempt to reload saved objects referenced by grid parameters (e.g. training frame, calibration frame), will fail if grid was saved without referenced objects.
Examples

```r
## Not run:
library(h2o)
h2o.init()

iris <- as.h2o(iris)

ntrees_opts = c(1, 5)
learn_rate_opts = c(0.1, 0.01)
size_of_hyper_space = length(ntrees_opts) * length(learn_rate_opts)

hyper_parameters = list(ntrees = ntrees_opts, learn_rate = learn_rate_opts)
# Tempdir is chosen arbitrarily. May be any valid folder on an H2O-supported filesystem.
baseline_grid <- h2o.grid("gbm", grid_id="gbm_grid_test", x=1:4, y=5, training_frame=iris,
hyper_params = hyper_parameters, export_checkpoints_dir = tempdir())
# Remove everything from the cluster or restart it
h2o.removeAll()
grid <- h2o.loadGrid(paste0(tempdir(), "/", baseline_grid@grid_id))

## End(Not run)
```

---

**h2o.loadModel**

*Load H2O Model from HDFS or Local Disk*

---

**Description**

Load a saved H2O model from disk. (Note that ensemble binary models can now be loaded using this method.)

**Usage**

```r
h2o.loadModel(path)
```

**Arguments**

- **path**: The path of the H2O Model to be imported.

**Value**

Returns a **H2OModel** object of the class corresponding to the type of model loaded.

**See Also**

- h2o.saveModel, H2OModel
Examples

```r
## Not run:
# library(h2o)
# h2o.init()
# prostate_path = system.file("extdata", "prostate.csv", package = "h2o")
# prostate = h2o.importFile(path = prostate_path)
# prostate_glm = h2o.glm(y = "CAPSULE", x = c("AGE", "RACE", "PSA", "DCAPS"),
# training_frame = prostate, family = "binomial", alpha = 0.5)
# glmmodel_path = h2o.saveModel(prostate_glm, dir = "/Users/UserName/Desktop")
# glmmodel_load = h2o.loadModel(glmmodel_path)

## End(Not run)
```

---

**h2o.load_frame**  
Load frame previously stored in H2O’s native format.

### Description

Load frame previously stored in H2O’s native format.

### Usage

```r
h2o.load_frame(frame_id, dir, force = TRUE)
```

### Arguments

- `frame_id`  
  the frame ID of the original frame
- `dir`  
  a filesystem location where to look for frame data
- `force`  
  logical. overwrite an already existing frame (defaults to true)

### Examples

```r
## Not run:
library(h2o)
h2o.init()

prostate_path = system.file("extdata", "prostate.csv", package = "h2o")
prostate = h2o.importFile(path = prostate_path)
h2o.save_frame(prostate, "/tmp/prostate")
prostate.key <- h2o.getId(prostate)
h2o.rm(prostate)
prostate <- h2o.load_frame(prostate.key, "/tmp/prostate")

## End(Not run)
```
h2o.log

Compute the logarithm of x

Description
Compute the logarithm of x

Usage
h2o.log(x)

Arguments
x An H2OFrame object.

See Also
Log for the base R implementation, log.

Examples
## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)
h2o.log(frame)

## End(Not run)

---

h2o.log10

Compute the log10 of x

Description
Compute the log10 of x

Usage
h2o.log10(x)

Arguments
x An H2OFrame object.
See Also

Log for the base R implementation, log10().

Examples

```r
## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)
h2o.log1p(frame)

## End(Not run)
```

---

h2o.log1p

Compute the log1p of x

Description

Compute the log1p of x

Usage

h2o.log1p(x)

Arguments

x

An H2OFrame object.

See Also

Log for the base R implementation, log1p().

Examples

```r
## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)
h2o.log1p(frame)

## End(Not run)
```
h2o.log2

Description

Compute the log2 of x

Usage

h2o.log2(x)

Arguments

x

An H2OFrame object.

See Also

Log for the base R implementation, log2()

Examples

## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)

h2o.log2(frame)

## End(Not run)

h2o.logAndEcho

Log a message on the server-side logs

Description

This is helpful when running several pieces of work one after the other on a single H2O cluster and you want to make a notation in the H2O server side log where one piece of work ends and the next piece of work begins.

Usage

h2o.logAndEcho(message)
h2o.logloss

Arguments

message A character string with the message to write to the log.

Details

h2o.logAndEcho sends a message to H2O for logging. Generally used for debugging purposes.

Description

Retrieves the log loss output for a H2OBinomialMetrics or H2OMultinomialMetrics object. If "train", "valid", and "xval" parameters are FALSE (default), then the training Log Loss value is returned. If more than one parameter is set to TRUE, then a named vector of Log Losses are returned, where the names are "train", "valid" or "xval".

Usage

h2o.logloss(object, train = FALSE, valid = FALSE, xval = FALSE)

Arguments

object a H2OModelMetrics object of the correct type.
train Retrieve the training Log Loss
valid Retrieve the validation Log Loss
xval Retrieve the cross-validation Log Loss

Examples

## Not run:
library(h2o)
h2o.init()

cars <- h2o.importFile(f)
cars["economy_20mpg"] <- as.factor(cars["economy_20mpg"])
predictors <- c("displacement", "power", "weight", "acceleration", "year")
response <- "economy_20mpg"
cars_splits <- h2o.splitFrame(data = cars, ratios = .8, seed = 1234)
train <- cars_splits[[1]]
valid <- cars_splits[[2]]
car_drf <- h2o.randomForest(x = predictors,
                           y = response,
                           training_frame = train,
                           validation_frame = valid)
h2o.logloss(car_drf, train = TRUE, valid = TRUE)

## End(Not run)
**h2o.ls**  
*List Keys on an H2O Cluster*

**Description**

Accesses a list of object keys in the running instance of H2O.

**Usage**

`h2o.ls()`

**Value**

Returns a list of hex keys in the current H2O instance.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
h2o.ls()
## End(Not run)
```

**h2o.lstrip**  
*Strip set from left*

**Description**

Return a copy of the target column with leading characters removed. The set argument is a string specifying the set of characters to be removed. If omitted, the set argument defaults to removing whitespace.

**Usage**

`h2o.lstrip(x, set = " ")`

**Arguments**

- `x` The column whose strings should be lstrip-ed.
- `set` string of characters to be removed
## Examples

```r
## Not run:
library(h2o)
str <- h2o("1234567890")
str <- str %>% lstrip("123") # Remove "123"

## End(Not run)
```

---

<h2>h2o.mae</h2>

<h3>Retrieve the Mean Absolute Error Value</h3>

### Description

Retrieves the mean absolute error (MAE) value from an H2O model. If "train", "valid", and "xval" parameters are FALSE (default), then the training MAE value is returned. If more than one parameter is set to TRUE, then a named vector of MAEs are returned, where the names are "train", "valid" or "xval".

### Usage

```r
h2o.mae(object, train = FALSE, valid = FALSE, xval = FALSE)
```

### Arguments

- **object**: An <b>H2OModel</b> object.
- **train**: Retrieve the training MAE.
- **valid**: Retrieve the validation set MAE if a validation set was passed in during model build time.
- **xval**: Retrieve the cross-validation MAE.

### Examples

```r
## Not run:
library(h2o)

h <- h2o.init()
fr <- as.h2o(iris)
m <- h2o.deeplearning(x = 2:5, y = 1, training_frame = fr)

h2o.mae(m)

## End(Not run)
```
**h2o.makeGLMModel**  
*Set betas of an existing H2O GLM Model*

**Description**
This function allows setting betas of an existing glm model.

**Usage**

```r
h2o.makeGLMModel(model, beta)
```

**Arguments**
- `model` an H2OModel corresponding from a h2o.glm call.
- `beta` a new set of betas (a named vector)

---

**h2o.make_metrics**  
*Create Model Metrics from predicted and actual values in H2O*

**Description**
Given predicted values (target for regression, class-1 probabilities or binomial or per-class probabilities for multinomial), compute a model metrics object.

**Usage**

```r
h2o.make_metrics(
  predicted,
  actuals,
  domain = NULL,
  distribution = NULL,
  weights = NULL,
  treatment = NULL,
  auc_type = "NONE",
  auuc_type = "AUTO",
  auuc_nbins = -1
)
```

**Arguments**
- `predicted` An H2OFrame containing predictions
- `actuals` An H2OFrame containing actual values
- `domain` Vector with response factors for classification.
- `distribution` Distribution for regression.
weights (optional) An H2OFrame containing observation weights.
treatment (optional, for uplift models only) An H2OFrame containing treatment column for uplift classification.

auc_type (optional) For multinomial classification you have to specify which type of aggregated AUC/AUCPR will be used to calculate this metric.

auuc_type (optional) For uplift binomial classification you have to specify which type of AUUC will be used to calculate this metric. Possibilities are gini, lift, gain, AUTO. Default is AUTO which means qini.

auuc_nbins (optional) For uplift binomial classification you have to specify number of bins to be used for calculation the AUUC. Default is -1, which means 1000.

Value

Returns an object of the H2OModelMetrics subclass.

Examples

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
prostate$CAPSULE <- as.factor(prostate$CAPSULE)
prostate_gbm <- h2o.gbm(3:9, "CAPSULE", prostate)
pred <- h2o.predict(prostate_gbm, prostate)[, 3] ## class-1 probability
h2o.make_metrics(pred, prostate$CAPSULE)
```

h2o.match

Value Matching in H2O

Description

match and %in% return values similar to the base R generic functions.

Usage

```r
h2o.match(x, table, nomatch = 0, incomparables = NULL)
match.H2OFrame(x, table, nomatch = 0, incomparables = NULL)
x %in% table
```
h2o.max

Returns the maxima of the input values.

**Arguments**

- `x`: An H2OFrame object.
- `na.rm`: logical, indicating whether missing values should be removed.

**See Also**

- `Extremes` for the base R implementation, `max()`.
Examples

```r
## Not run:
library(h2o)
h2o.init()

iris <- h2o.importFile(f)
h2o.max(iris["petal_len"], na.rm = TRUE)

## End(Not run)
```

---

**h2o.mean**

*Compute the frame’s mean by-column (or by-row).*

### Description

Compute the frame’s mean by-column (or by-row).

### Usage

```r
h2o.mean(x, na.rm = FALSE, axis = 0, return_frame = FALSE, ...)
```

#### S3 method for class 'H2OFrame'

```r
mean(x, na.rm = FALSE, axis = 0, return_frame = FALSE, ...)
```

### Arguments

- **x**: An H2OFrame object.
- **na.rm**: logical. Indicate whether missing values should be removed.
- **axis**: integer. Indicate whether to calculate the mean down a column (0) or across a row (1). NOTE: This is only applied when return_frame is set to TRUE. Otherwise, this parameter is ignored.
- **return_frame**: logical. Indicate whether to return an H2O frame or a list. Default is FALSE (returns a list).
- **...**: Further arguments to be passed from or to other methods.

### Value

Returns a list containing the mean for each column (NaN for non-numeric columns) if return_frame is set to FALSE. If return_frame is set to TRUE, then it will return an H2O frame with means per column or row (depends on axis argument).

### See Also

`Round` for base R implementation, `mean()` and `colSums` for the base R implementation, `colMeans()`.
## Examples

```r
## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
# Default behavior. Will return list of means per column.
h2o.mean(prostate$AGE)
# return_frame set to TRUE. This will return an H2O Frame
# with mean per row or column (depends on axis argument)
h2o.mean(prostate, na.rm = TRUE, axis = 1, return_frame = TRUE)

## End(Not run)
```

---

### h2o.mean_per_class_error

*Retrieve the mean per class error*

**Description**

Retrieves the mean per class error from an `H2OBinomialMetrics`. If "train", "valid", and "xval" parameters are FALSE (default), then the training mean per class error value is returned. If more than one parameter is set to TRUE, then a named vector of mean per class errors are returned, where the names are "train", "valid" or "xval".

**Usage**

```r
h2o.mean_per_class_error(object, train = FALSE, valid = FALSE, xval = FALSE)
```

**Arguments**

- `object`: An `H2OBinomialMetrics` object.
- `train`: Retrieve the training mean per class error
- `valid`: Retrieve the validation mean per class error
- `xval`: Retrieve the cross-validation mean per class error

**See Also**

- `h2o.mse` for MSE, and `h2o.metric` for the various threshold metrics. See `h2o.performance` for creating `H2OModelMetrics` objects.
Examples

```r
## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(prostate_path)

prostate[, 2] <- as.factor(prostate[, 2])
model <- h2o.gbm(x = 3:9, y = 2, training_frame = prostate, distribution = "bernoulli")
perf <- h2o.performance(model, prostate)
h2o.mean_per_class_error(perf)
h2o.mean_per_class_error(model, train=TRUE)

## End(Not run)
```

---

**h2o.mean_residual_deviance**

*Retrieve the Mean Residual Deviance value*

**Description**

Retrieves the Mean Residual Deviance value from an H2O model. If "train", "valid", and "xval" parameters are FALSE (default), then the training Mean Residual Deviance value is returned. If more than one parameter is set to TRUE, then a named vector of Mean Residual Deviances are returned, where the names are "train", "valid" or "xval".

**Usage**

```r
h2o.mean_residual_deviance(object, train = FALSE, valid = FALSE, xval = FALSE)
```

**Arguments**

- **object**: An H2OModel object.
- **train**: Retrieve the training Mean Residual Deviance
- **valid**: Retrieve the validation Mean Residual Deviance
- **xval**: Retrieve the cross-validation Mean Residual Deviance

**Examples**

```r
## Not run:
library(h2o)

h <- h2o.init()
fr <- as.h2o(iris)
m <- h2o.deeplearning(x = 2:5, y = 1, training_frame = fr)
```
h2o.median

Description

Compute the median of an H2OFrame.

Usage

h2o.median(x, na.rm = TRUE)

## S3 method for class 'H2OFrame'
median(x, na.rm = TRUE)

Arguments

x
An H2OFrame object.

na.rm
a logical, indicating whether na’s are omitted.

Value

Returns a list containing the median for each column (NaN for non-numeric columns)

Examples

## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
h2o.median(prostate)

## End(Not run)
h2o.melt

Converts a frame to key-value representation while optionally skipping NA values. Inverse operation to h2o.pivot.

Description

Pivot the frame designated by the three columns: index, column, and value. Index and column should be of type enum, int, or time. For cases of multiple indexes for a column label, the aggregation method is to pick the first occurrence in the data frame.

Usage

h2o.melt(
  x,
  id_vars,
  value_vars = NULL,
  var_name = "variable",
  value_name = "value",
  skipna = FALSE
)

Arguments

- **x**: an H2OFrame
- **id_vars**: the columns used as identifiers
- **value_vars**: what columns will be converted to key-value pairs (optional, if not specified complement to id_vars will be used)
- **var_name**: name of the key-column (default: "variable")
- **value_name**: name of the value-column (default: "value")
- **skipna**: if enabled, do not include NAs in the result (default: FALSE)

Value

an unpivoted H2OFrame
**Description**

Merges two H2OFrame objects with the same arguments and meanings as merge() in base R. However, we do not support all=TRUE, all.x=TRUE and all.y=TRUE. The default method is auto and it will default to the radix method. The radix method will return the correct merge result regardless of duplicated rows in the right frame. In addition, the radix method can perform merge even if you have string columns in your frames. If there are duplicated rows in your right frame, they will not be included if you use the hash method. The hash method cannot perform merge if you have string columns in your left frame. Hence, we consider the radix method superior to the hash method and is the default method to use.

**Usage**

```r
h2o.merge(
  x, y,
  by = intersect(names(x), names(y)),
  by.x = by,
  by.y = by,
  all = FALSE,
  all.x = all,
  all.y = all,
  method = "auto"
)
```

**Arguments**

- `x, y` H2OFrame objects
- `by` columns used for merging by default the common names
- `by.x` x columns used for merging by name or number
- `by.y` y columns used for merging by name or number
- `all` TRUE includes all rows in x and all rows in y even if there is no match to the other
- `all.x` If all.x is true, all rows in the x will be included, even if there is no matching row in y, and vice-versa for all.y.
- `all.y` see `all.x`
- `method` auto(default), radix, hash

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
left <- data.frame(fruit = c("apple", "orange", "banana", "lemon", "strawberry", "blueberry"),
  color <- c("red", "orange", "yellow", "yellow", "red", "blue"))
right <- data.frame(fruit = c("apple", "orange", "banana", "lemon", "strawberry", "watermelon"),
  citrus <- c(FALSE, TRUE, FALSE, TRUE, FALSE, FALSE))
left_hf <- as.h2o(left)
```
right_hf <- as.h2o(right)
merged <- h2o.merge(left_hf, right_hf, all.x = TRUE)

## End(Not run)

---

<h20.metric>  
**H2O Model Metric Accessor Functions**

**Description**
A series of functions that retrieve model metric details.

**Usage**

- `h2o.metric(object, thresholds, metric, transform = NULL)`
- `h2o.F0point5(object, thresholds)`
- `h2o.F1(object, thresholds)`
- `h2o.F2(object, thresholds)`
- `h2o.accuracy(object, thresholds)`
- `h2o.error(object, thresholds)`
- `h2o.maxPerClassError(object, thresholds)`
- `h2o.mean_per_class_accuracy(object, thresholds)`
- `h2o.mcc(object, thresholds)`
- `h2o.precision(object, thresholds)`
- `h2o.tpr(object, thresholds)`
- `h2o.fpr(object, thresholds)`
- `h2o.fnr(object, thresholds)`
- `h2o.tnr(object, thresholds)`
- `h2o.recall(object, thresholds)`
- `h2o.sensitivity(object, thresholds)`
- `h2o.fallout(object, thresholds)`
h2o.missrate(object, thresholds)

h2o.specificity(object, thresholds)

**Arguments**

- **object**: An `H2OModelMetrics` object of the correct type.
- **thresholds**: (Optional) A value or a list of values between 0.0 and 1.0. If not set, then all thresholds will be returned. If "max", then the threshold maximizing the metric will be used.
- **metric**: (Optional) the metric to retrieve. If not set, then all metrics will be returned.
- **transform**: (Optional) a list describing a transformer for the given metric, if any. e.g. `transform=list(op=foo_fn, name="foo")` will rename the given metric to "foo" and apply function `foo_fn` to the metric values.

**Details**

Many of these functions have an optional thresholds parameter. Currently only increments of 0.1 are allowed. If not specified, the functions will return all possible values. Otherwise, the function will return the value for the indicated threshold.

Currently, the these functions are only supported by `H2OBinomialMetrics` objects.

**Value**

Returns either a single value, or a list of values.

**See Also**

- `h2o.auc` for AUC, `h2o.giniCoef` for the GINI coefficient, and `h2o.mse` for MSE. See `h2o.performance` for creating `H2OModelMetrics` objects.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(prostate_path)
prostate$CAPSULE <- as.factor(prostate$CAPSULE)
model <- h2o.gbm(x = 3:9, y = 2, training_frame = prostate, distribution = "bernoulli")
perf <- h2o.performance(model, prostate)
h2o.F1(perf)
## End(Not run)
```
**h2o.min**

*Returns the minima of the input values.*

**Description**

Returns the minima of the input values.

**Usage**

h2o.min(x, na.rm = FALSE)

**Arguments**

- **x**
  - An H2OFrame object.
- **na.rm**
  - logical. indicating whether missing values should be removed.

**See Also**

`Extremes` for the base R implementation, `min()`.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

iris <- h2o.importFile(f)
h2o.min(iris["sepal_len"], na.rm = TRUE)

## End(Not run)
```

---

**h2o.mktime**

*Compute msec since the Unix Epoch*

**Description**

Compute msec since the Unix Epoch
Usage

```
    h2o.mktime(
            year = 1970,
            month = 0,
            day = 0,
            hour = 0,
            minute = 0,
            second = 0,
            msec = 0
        )
```

Arguments

- **year**
  - Defaults to 1970
- **month**
  - zero based (months are 0 to 11)
- **day**
  - zero based (days are 0 to 30)
- **hour**
- **minute**
- **second**
- **msec**

Examples

```r
## Not run:
library(h2o)
h2o.init()

x = as.h2o(c(2018, 3, 2, 6, 32, 0, 0))
h2o.mktime(x)
## End(Not run)
```

---

**h2o.modelSelection**

H2O ModelSelection is used to build the best model with one predictor, two predictors, ... up to max_predictor_number specified in the algorithm parameters when mode=allsubsets. The best model is the one with the highest R2 value. When mode=maxr, the model returned is no longer guaranteed to have the best R2 value.

---

**Description**

H2O ModelSelection is used to build the best model with one predictor, two predictors, ... up to max_predictor_number specified in the algorithm parameters when mode=allsubsets. The best model is the one with the highest R2 value. When mode=maxr, the model returned is no longer guaranteed to have the best R2 value.
Usage

h2o.modelSelection(
  x,
  y,
  training_frame,
  model_id = NULL,
  validation_frame = NULL,
  nfolds = 0,
  seed = -1,
  fold_assignment = c("AUTO", "Random", "Modulo", "Stratified"),
  fold_column = NULL,
  ignore_constcols = TRUE,
  score_each_iteration = FALSE,
  score_iteration_interval = 0,
  offset_column = NULL,
  weights_column = NULL,
  family = c("AUTO", "gaussian", "binomial", "fractionalbinomial", "quasibinomial",
    "poisson", "gamma", "tweedie", "negativebinomial"),
  link = c("family_default", "identity", "logit", "log", "inverse", "tweedie",
    "ologit"),
  tweedie_variance_power = 0,
  tweedie_link_power = 0,
  theta = 0,
  solver = c("AUTO", "IRLSM", "L_BFGS", "COORDINATE_DESCENT_NAIVE",
    "COORDINATE_DESCENT", "GRADIENT_DESCENT_LH", "GRADIENT_DESCENT_SQERR"),
  alpha = NULL,
  lambda = NULL,
  lambda_search = FALSE,
  early_stopping = FALSE,
  nlambdas = 0,
  standardize = TRUE,
  missing_values_handling = c("MeanImputation", "Skip", "PlugValues"),
  plug_values = NULL,
  compute_p_values = FALSE,
  remove_collinear_columns = FALSE,
  intercept = FALSE,
  non_negative = FALSE,
  max_iterations = 0,
  objective_epsilon = 0,
  beta_epsilon = 0,
  gradient_epsilon = 0,
  startval = NULL,
  prior = 0,
  cold_start = FALSE,
  lambda_min_ratio = 0,
  beta_constraints = NULL,
  max_active_predictors = -1,
  obj_reg = 0,
stopping_rounds = 0,
stopping_metric = c("AUTO", "deviance", "logloss", "MSE", "RMSE", "MAE", "RMSLE", "AUC", "AUCPR", "lift_top_group", "misclassification", "mean_per_class_error", "custom", "custom_increasing"),
stopping_tolerance = 0.001,
balance_classes = FALSE,
class_sampling_factors = NULL,
max_after_balance_size = 5,
max_runtime_secs = 0,
custom_metric_func = NULL,
nparallelism = 0,
max_predictor_number = 1,
min_predictor_number = 1,
mode = c("allsubsets", "maxr", "backward"),
p_values_threshold = 0
)

Arguments

x (Optional) A vector containing the names or indices of the predictor variables to use in building the model. If x is missing, then all columns except y are used.

y The name or column index of the response variable in the data. The response must be either a numeric or a categorical/factor variable. If the response is numeric, then a regression model will be trained, otherwise it will train a classification model.

training_frame Id of the training data frame.
model_id Destination id for this model; auto-generated if not specified.
validation_frame Id of the validation data frame.

nfolds Number of folds for K-fold cross-validation (0 to disable or >= 2). Defaults to 0.

seed Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).

fold_assignment Cross-validation fold assignment scheme, if fold_column is not specified. The 'Stratified' option will stratify the folds based on the response variable, for classification problems. Must be one of: "AUTO", "Random", "Modulo", "Stratified". Defaults to AUTO.

fold_column Column with cross-validation fold index assignment per observation.
ignore_const_cols Logical. Ignore constant columns. Defaults to TRUE.

score_each_iteration Logical. Whether to score during each iteration of model training. Defaults to FALSE.
score_iteration_interval Perform scoring for every score_iteration_interval iterations Defaults to 0.
offset_column  Offset column. This will be added to the combination of columns before applying the link function.

weights_column  Column with observation weights. Giving some observation a weight of zero is equivalent to excluding it from the dataset; giving an observation a relative weight of 2 is equivalent to repeating that row twice. Negative weights are not allowed. Note: Weights are per-row observation weights and do not increase the size of the data frame. This is typically the number of times a row is repeated, but non-integer values are supported as well. During training, rows with higher weights matter more, due to the larger loss function pre-factor. If you set weight = 0 for a row, the returned prediction frame at that row is zero and this is incorrect. To get an accurate prediction, remove all rows with weight == 0.

family  Family. For MaxR, only gaussian. For backward, ordinal and multinomial families are not supported. Must be one of: "AUTO", "gaussian", "binomial", "fractionalbinomial", "quasibinomial", "poisson", "gamma", "tweedie", "negativebinomial". Defaults to AUTO.

link  Link function. Must be one of: "family_default", "identity", "logit", "log", "inverse", "tweedie", "ologit". Defaults to family_default.

tweedie_variance_power  Tweedie variance power Defaults to 0.

tweedie_link_power  Tweedie link power Defaults to 0.

theta  Theta Defaults to 0.

solver  AUTO will set the solver based on given data and the other parameters. IRLSM is fast on on problems with small number of predictors and for lambda-search with L1 penalty, L_BFGS scales better for datasets with many columns. Must be one of: "AUTO", "IRLSM", "L_BFGS", "COORDINATE_DESCENT_NAIVE", "COORDINATE_DESCENT", "GRADIENT_DESCENT_LH", "GRADIENT_DESCENT_SQERR". Defaults to IRLSM.

alpha  Distribution of regularization between the L1 (Lasso) and L2 (Ridge) penalties. A value of 1 for alpha represents Lasso regression, a value of 0 produces Ridge regression, and anything in between specifies the amount of mixing between the two. Default value of alpha is 0 when SOLVER = 'L-BFGS'; 0.5 otherwise.

lambda  Regularization strength

lambda_search  Logical. Use lambda search starting at lambda max, given lambda is then interpreted as lambda min. Defaults to FALSE.

early_stopping  Logical. Stop early when there is no more relative improvement on train or validation (if provided). Defaults to FALSE.

nlambdas  Number of lambdas to be used in a search. Default indicates: If alpha is zero, with lambda search set to True, the value of nlambdas is set to 30 (fewer lambdas are needed for ridge regression) otherwise it is set to 100. Defaults to 0.

standardize  Logical. Standardize numeric columns to have zero mean and unit variance. Defaults to TRUE.

missing_values_handling  Handling of missing values. Either MeanImputation, Skip or PlugValues. Must be one of: "MeanImputation", "Skip", "PlugValues". Defaults to MeanImputation.
h2o.modelSelection

plug_values
Plug Values (a single row frame containing values that will be used to impute missing values of the training/validation frame, use with conjunction missing_values_handling = PlugValues)

compute_p_values
Logical. Request p-values computation. p-values work only with IRLSM solver and no regularization Defaults to FALSE.

remove_collinear_columns
Logical. In case of linearly dependent columns, remove some of the dependent columns Defaults to FALSE.

intercept
Logical. Include constant term in the model Defaults to FALSE.

non_negative
Logical. Restrict coefficients (not intercept) to be non-negative Defaults to FALSE.

max_iterations
Maximum number of iterations Defaults to 0.

objective_epsilon
Converge if objective value changes less than this. Default indicates: If lambda_search is set to True the value of objective_epsilon is set to .0001. If the lambda_search is set to False and lambda is equal to zero, the value of objective_epsilon is set to .000001, for any other value of lambda the default value of objective_epsilon is set to .0001. Defaults to 0.

beta_epsilon
Converge if beta changes less (using L-infinity norm) than beta epsilon, ONLY applies to IRLSM solver Defaults to 0.

gradient_epsilon
Converge if objective changes less (using L-infinity norm) than this, ONLY applies to L-BFGS solver. Default indicates: If lambda_search is set to False and lambda is equal to zero, the default value of gradient_epsilon is equal to .000001, otherwise the default value is .0001. If lambda_search is set to True, the conditional values above are 1E-8 and 1E-6 respectively. Defaults to 0.

startval
double array to initialize fixed and random coefficients for HGLM, coefficients for GLM.

prior
Prior probability for y==1. To be used only for logistic regression iff the data has been sampled and the mean of response does not reflect reality. Defaults to 0.

cold_start
Logical. Only applicable to multiple alpha/lambda values. If false, build the next model for next set of alpha/lambda values starting from the values provided by current model. If true will start GLM model from scratch. Defaults to FALSE.

lambda_min_ratio
Minimum lambda used in lambda search, specified as a ratio of lambda_max (the smallest lambda that drives all coefficients to zero). Default indicates: if the number of observations is greater than the number of variables, then lambda_min_ratio is set to 0.0001; if the number of observations is less than the number of variables, then lambda_min_ratio is set to 0.01. Defaults to 0.

beta_constraints
Beta constraints
max_active_predictors
Maximum number of active predictors during computation. Use as a stopping criterion to prevent expensive model building with many predictors. Default indicates: If the IRLSM solver is used, the value of max_active_predictors is set to 5000 otherwise it is set to 100000000. Defaults to -1.

obj_reg
Likelihood divider in objective value computation, default is \(1/\text{nobs}\) Defaults to 0.

stopping_rounds
Early stopping based on convergence of stopping_metric. Stop if simple moving average of length \(k\) of the stopping_metric does not improve for \(k = \text{stopping_rounds}\) scoring events (0 to disable) Defaults to 0.

stopping_metric
Metric to use for early stopping (AUTO: logloss for classification, deviance for regression and anomaly_score for Isolation Forest). Note that custom and custom_increasing can only be used in GBM and DRF with the Python client. Must be one of: "AUTO", "deviance", "logloss", "MSE", "RMSE", "MAE", "RMSLE", "AUC", "AUCPR", "lift_top_group", "misclassification", "mean_per_class_error", "custom", "custom_increasing". Defaults to AUTO.

stopping_tolerance
Relative tolerance for metric-based stopping criterion (stop if relative improvement is not at least this much) Defaults to 0.001.

balance_classes
Logical. Balance training data class counts via over/under-sampling (for imbalanced data). Defaults to FALSE.

class_sampling_factors
Desired over/under-sampling ratios per class (in lexicographic order). If not specified, sampling factors will be automatically computed to obtain class balance during training. Requires balance_classes.

max_after_balance_size
Maximum relative size of the training data after balancing class counts (can be less than 1.0). Requires balance_classes. Defaults to 5.0.

max_runtime_secs
Maximum allowed runtime in seconds for model training. Use 0 to disable. Defaults to 0.

custom_metric_func
Reference to custom evaluation function, format: ‘language:KeyName=funcName’

nparallelism
Number of models to build in parallel. Defaults to 0.0 which is adaptive to the system capability Defaults to 0.

max_predictor_number
Maximum number of predictors to be considered when building GLM models. Defaults to 1. Defaults to 1.

min_predictor_number
For mode = 'backward' only. Minimum number of predictors to be considered when building GLM models starting with all predictors to be included. Defaults to 1. Defaults to 1.

mode
Mode: Used to choose model selection algorithms to use. Options include 'allsubsets' for all subsets, 'maxr' for MaxR, 'backward' for backward selection Must be one of: "allsubsets", "maxr", "backward". Defaults to maxr.
p_values_threshold
For mode='backward' only. If specified, will stop the model building process when all coefficients p-values drop below this threshold. Defaults to 0.

Examples

```r
## Not run:
library(h2o)
h2o.init()
# Run ModelSelection of VOL ~ all predictors
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
prostate$CAPSULE <- as.factor(prostate$CAPSULE)
model <- h2o.modelSelection(y="VOL", x=c("RACE","AGE","RACE","DPROS"), training_frame=prostate)
## End(Not run)
```

h2o.model_correlation  Model Prediction Correlation

Description
Get a data.frame containing the correlation between the predictions of the models. For classification, frequency of identical predictions is used. By default, models are ordered by their similarity (as computed by hierarchical clustering).

Usage

```
h2o.model_correlation(object, newdata, top_n = 20, cluster_models = TRUE)
```

Arguments

- **object**: A list of H2O models, an H2O AutoML instance, or an H2OFrame with a 'model_id' column (e.g. H2OAutoML leaderboard).
- **newdata**: An H2O Frame. Predictions from the models will be generated using this frame, so this should be a holdout set.
- **top_n**: (DEPRECATED) Integer specifying the number models shown in the heatmap (used only with an AutoML object, and based on the leaderboard ranking. Defaults to 20.
- **cluster_models**: Logical. Order models based on their similarity. Defaults to TRUE.

Value
A data.frame containing variable importance.
Examples

```r
## Not run:
library(h2o)
h2o.init()

# Import the wine dataset into H2O:
df <- h2o.importFile(f)

# Set the response
response <- "quality"

# Split the dataset into a train and test set:
splits <- h2o.splitFrame(df, ratios = 0.8, seed = 1)
train <- splits[[1]]
test <- splits[[2]]

# Build and train the model:
aml <- h2o.automl(y = response,
                  training_frame = train,
                  max_models = 10,
                  seed = 1)

# Create the model correlation
model_correlation <- h2o.model_correlation(aml, test)
print(model_correlation)

## End(Not run)
```

---

**h2o.model_correlation_heatmap**

*Model Prediction Correlation Heatmap*

**Description**

This plot shows the correlation between the predictions of the models. For classification, frequency of identical predictions is used. By default, models are ordered by their similarity (as computed by hierarchical clustering).

**Usage**

```r
h2o.model_correlation_heatmap(
  object,
  newdata,
  top_n = 20,
  cluster_models = TRUE,
  triangular = TRUE
)
```
Arguments

object  A list of H2O models, an H2O AutoML instance, or an H2OFrame with a 'model_id' column (e.g. H2OAutoML leaderboard).

newdata  An H2O Frame. Predictions from the models will be generated using this frame, so this should be a holdout set.

top_n  Integer specifying the number models shown in the heatmap (used only with an AutoML object, and based on the leaderboard ranking. Defaults to 20.

cluster_models  Logical. Order models based on their similarity. Defaults to TRUE.

triangular  Print just the lower triangular part of correlation matrix. Defaults to TRUE.

Value

A ggplot2 object.

Examples

```r
## Not run:
library(h2o)
h2o.init()

# Import the wine dataset into H2O:
df <- h2o.importFile(f)

# Set the response
response <- "quality"

# Split the dataset into a train and test set:
splits <- h2o.splitFrame(df, ratios = 0.8, seed = 1)
train <- splits[[1]]
test <- splits[[2]]

# Build and train the model:
aml <- h2o.automl(y = response,
training_frame = train,
max_models = 10,
seed = 1)

# Create the model correlation heatmap
model_correlation_heatmap <- h2o.model_correlation_heatmap(aml, test)
print(model_correlation_heatmap)
```

## End(Not run)
Description

Provides the method `h2o.mojo_predict_csv` with which you can predict a MOJO model from R.

Usage

```r
h2o.mojo_predict_csv(
  input_csv_path,
  mojo_zip_path,
  output_csv_path = NULL,
  genmodel_jar_path = NULL,
  classpath = NULL,
  java_options = NULL,
  verbose = F,
  setInvNumNA = F
)
```

Arguments

- **input_csv_path**: Path to input CSV file.
- **mojo_zip_path**: Path to MOJO zip downloaded from H2O.
- **output_csv_path**: Optional, path to the output CSV file with computed predictions. If NULL (default), then predictions will be saved as prediction.csv in the same folder as the MOJO zip.
- **genmodel_jar_path**: Optional, path to genmodel jar file. If NULL (default) then the h2o-genmodel.jar in the same folder as the MOJO zip will be used.
- **classpath**: Optional, specifies custom user defined classpath which will be used when scoring. If NULL (default) then the default classpath for this MOJO model will be used.
- **java_options**: Optional, custom user defined options for Java. By default `-Xmx4g -XX:ReservedCodeCacheSize=256m` is used.
- **verbose**: Optional, if TRUE, then additional debug information will be printed. FALSE by default.
- **setInvNumNA**: Optional, if TRUE, then then for an string that cannot be parsed into a number an N/A value will be produced, if false the command will fail. FALSE by default.

Value

Returns a data.frame containing computed predictions
Description

Provides the method `h2o.mojo_predict_df` with which you can predict a MOJO model from R.

Usage

```r
h2o.mojo_predict_df(
  frame,  
  mojo_zip_path,  
  genmodel_jar_path = NULL,  
  classpath = NULL,  
  java_options = NULL,  
  verbose = F,  
  setInvNumNA = F
)
```

Arguments

- `frame` : data.frame to score.
- `mojo_zip_path` : Path to MOJO zip downloaded from H2O.
- `genmodel_jar_path` : Optional, path to genmodel jar file. If NULL (default) then the `h2o-genmodel.jar` in the same folder as the MOJO zip will be used.
- `classpath` : Optional, specifies custom user defined classpath which will be used when scoring. If NULL (default) then the default classpath for this MOJO model will be used.
- `java_options` : Optional, custom user defined options for Java. By default `-Xmx4g -XX:ReservedCodeCacheSize=256m` is used.
- `verbose` : Optional, if TRUE, then additional debug information will be printed. FALSE by default.
- `setInvNumNA` : Optional, if TRUE, then for an string that cannot be parsed into a number an N/A value will be produced, if false the command will fail. FALSE by default.

Value

Returns a data.frame containing computed predictions.
**h2o.month**

*Convert Milliseconds to Months in H2O Datasets*

**Description**

Converts the entries of an H2OFrame object from milliseconds to months (on a 1 to 12 scale).

**Usage**

- `h2o.month(x)`
- `month(x)`

**Arguments**

- `x` An H2OFrame object.

**Value**

An H2OFrame object containing the entries of `x` converted to months of the year.

**See Also**

- `h2o.year`

---

**h2o.mse**

*Retrieves Mean Squared Error Value*

**Description**

Retrieves the mean squared error value from an H2OModelMetrics object. If "train", "valid", and "xval" parameters are FALSE (default), then the training MSE value is returned. If more than one parameter is set to TRUE, then a named vector of MSESs are returned, where the names are "train", "valid" or "xval".

**Usage**

- `h2o.mse(object, train = FALSE, valid = FALSE, xval = FALSE)`.  

---
Arguments

- **object**: An H2OModelMetrics object of the correct type.
- **train**: Retrieve the training MSE.
- **valid**: Retrieve the validation MSE.
- **xval**: Retrieve the cross-validation MSE.

Details

This function only supports H2OBinomialMetrics, H2OMultinomialMetrics, and H2ORegressionMetrics objects.

See Also

- h2o.auc for AUC,
- h2o.mse for MSE,
- h2o.metric for the various threshold metrics.
- See h2o.performance for creating H2OModelMetrics objects.

Examples

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(prostate_path)
prostate[, 2] <- as.factor(prostate[, 2])
model <- h2o.gbm(x = 3:9, y = 2, training_frame = prostate, distribution = "bernoulli")
perf <- h2o.performance(model, prostate)
h2o.mse(perf)
## End(Not run)
```

**h2o.multinomial_aucpr_table**

Retrieve the all PR AUC values in a table (One to Rest, One to One, macro and weighted average) for multinomial classification.

Description

Retrieves the PR AUC table from an H2OMultinomialMetrics. If "train", "valid", and "xval" parameters are FALSE (default), then the training PR AUC table is returned. If more than one parameter is set to TRUE, then a named vector of PR AUC tables are returned, where the names are "train", "valid" or "xval".

Usage

```r
h2o.multinomial_aucpr_table(object, train = FALSE, valid = FALSE, xval = FALSE)
```
Arguments

- **object**: An `H2OMultinomialMetrics` object.
- **train**: Retrieve the training PR AUC table
- **valid**: Retrieve the validation PR AUC table
- **xval**: Retrieve the cross-validation PR AUC table

See Also

- `h2o.giniCoef` for the Gini coefficient, `h2o.mse` for MSE, and `h2o.metric` for the various threshold metrics. See `h2o.performance` for creating H2OModelMetrics objects.

Examples

```r
## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(prostate_path)

prostate[, 2] <- as.factor(prostate[, 2])
model <- h2o.gbm(x = 3:9, y = 2, training_frame = prostate, distribution = "bernoulli")
perf <- h2o.performance(model, prostate)
h2o.multinomial_auc_table(perf)

## End(Not run)
```

**h2o.multinomial_auc_table**

*Retrieve the all AUC values in a table (One to Rest, One to One, macro and weighted average) for mutlinomial classification.*

Description

Retrieves the AUC table from an `H2OMultinomialMetrics`. If "train", "valid", and "xval" parameters are FALSE (default), then the training AUC table is returned. If more than one parameter is set to TRUE, then a named vector of AUC tables are returned, where the names are "train", "valid" or "xval".

Usage

`h2o.multinomial_auc_table(object, train = FALSE, valid = FALSE, xval = FALSE)`

Arguments

- **object**: An `H2OMultinomialMetrics` object.
- **train**: Retrieve the training AUC table
- **valid**: Retrieve the validation AUC table
- **xval**: Retrieve the cross-validation AUC table
See Also

`h2o.giniCoef` for the Gini coefficient, `h2o.mse` for MSE, and `h2o.metric` for the various threshold metrics. See `h2o.performance` for creating H2OModelMetrics objects.

Examples

```r
## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(prostate_path)

prostate[, 2] <- as.factor(prostate[, 2])
model <- h2o.gbm(x = 3:9, y = 2, training_frame = prostate, distribution = "bernoulli")
perf <- h2o.performance(model, prostate)
h2o.multinomial_auc_table(perf)

## End(Not run)
```

---

**h2o.nacnt**

*Count of NAs per column*

Description

Gives the count of NAs per column.

Usage

```r
h2o.nacnt(x)
```

Arguments

- `x` An H2OFrame object.

Value

Returns a list containing the count of NAs per column.

Examples

```r
## Not run:
library(h2o)
h2o.init()

iris_hf <- as.h2o(iris)
h2o.nacnt(iris_hf) # should return all 0s
h2o.insertMissingValues(iris_hf)
h2o.nacnt(iris_hf)
```
h2o.naiveBayes

**Compute naive Bayes probabilities on an H2O dataset.**

**Description**

The naive Bayes classifier assumes independence between predictor variables conditional on the response, and a Gaussian distribution of numeric predictors with mean and standard deviation computed from the training dataset. When building a naive Bayes classifier, every row in the training dataset that contains at least one NA will be skipped completely. If the test dataset has missing values, then those predictors are omitted in the probability calculation during prediction.

**Usage**

```r
h2o.naiveBayes(
  x,
  y,
  training_frame,
  model_id = NULL,
  nfolds = 0,
  seed = -1,
  fold_assignment = c("AUTO", "Random", "Modulo", "Stratified"),
  fold_column = NULL,
  keep_cross_validation_models = TRUE,
  keep_cross_validation_predictions = FALSE,
  keep_cross_validation_fold_assignment = FALSE,
  validation_frame = NULL,
  ignore_const_cols = TRUE,
  score_each_iteration = FALSE,
  balance_classes = FALSE,
  class_sampling_factors = NULL,
  max_after_balance_size = 5,
  laplace = 0,
  threshold = 0.001,
  min_sdev = 0.001,
  eps = 0,
  eps_sdev = 0,
  min_prob = 0.001,
  eps_prob = 0,
  compute_metrics = TRUE,
  max_runtime_secs = 0,
  export_checkpoints_dir = NULL,
  gainslift_bins = -1,
  auc_type = c("AUTO", "NONE", "MACRO_OVR", "WEIGHTED_OVR", "MACRO_OVO",
               "WEIGHTED_OVO")
)
```
Arguments

- **x** (Optional) A vector containing the names or indices of the predictor variables to use in building the model. If x is missing, then all columns except y are used.
- **y** The name or column index of the response variable in the data. The response must be either a numeric or a categorical/factor variable. If the response is numeric, then a regression model will be trained, otherwise it will train a classification model.
- **training_frame** Id of the training data frame.
- **model_id** Destination id for this model; auto-generated if not specified.
- **nfolds** Number of folds for K-fold cross-validation (0 to disable or >= 2). Defaults to 0.
- **seed** Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).
- **fold_assignment** Cross-validation fold assignment scheme, if fold_column is not specified. The 'Stratified' option will stratify the folds based on the response variable, for classification problems. Must be one of: "AUTO", "Random", "Modulo", "Stratified". Defaults to AUTO.
- **fold_column** Column with cross-validation fold index assignment per observation.
- **keep_cross_validation_models** Logical. Whether to keep the cross-validation models. Defaults to TRUE.
- **keep_cross_validation_predictions** Logical. Whether to keep the predictions of the cross-validation models. Defaults to FALSE.
- **keep_cross_validation_fold_assignment** Logical. Whether to keep the cross-validation fold assignment. Defaults to FALSE.
- **validation_frame** Id of the validation data frame.
- **ignore_const_cols** Logical. Ignore constant columns. Defaults to TRUE.
- **score_each_iteration** Logical. Whether to score during each iteration of model training. Defaults to FALSE.
- **balance_classes** Logical. Balance training data class counts via over/under-sampling (for imbalanced data). Defaults to FALSE.
- **class_sampling_factors** Desired over/under-sampling ratios per class (in lexicographic order). If not specified, sampling factors will be automatically computed to obtain class balance during training. Requires balance_classes.
- **max_after_balance_size** Maximum relative size of the training data after balancing class counts (can be less than 1.0). Requires balance_classes. Defaults to 5.0.
h2o.names

Column names of an H2OFrame

Description

Column names of an H2OFrame

### Examples

```r
## Not run:
# h2o.init()
votes_path <- system.file("extdata", "housevotes.csv", package = "h2o")
votes <- h2o.uploadFile(path = votes_path, header = TRUE)
h2o.naiveBayes(x = 2:17, y = 1, training_frame = votes, laplace = 3)

## End(Not run)
```
Usage

h2o.names(x)

Arguments

x An H2OFrame object.

See Also

names for the base R implementation.

Examples

## Not run:
library(h2o)
h2o.init()

iris <- h2o.importFile(f)
h2o.names(iris)

## End(Not run)

---

h2o.na_omit Remove Rows With NAs

Description

Remove Rows With NAs

Usage

h2o.na_omit(object, ...)

Arguments

object H2OFrame object
...
... Ignored

Value

Returns an H2OFrame object containing non-NA rows.
Examples

```r
## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)
h2o.na_omit(frame)
## End(Not run)
```

---

<table>
<thead>
<tr>
<th>h2o.nchar</th>
<th>String length</th>
</tr>
</thead>
</table>

Description

String length

Usage

```r
h2o.nchar(x)
```

Arguments

- `x` The column whose string lengths will be returned.

Examples

```r
## Not run:
library(h2o)
h2o.init()
string_to_nchar <- as.h2o("r tutorial")
nchar_string <- h2o.nchar(string_to_nchar)
## End(Not run)
```
h2o.ncol

Return the number of columns present in x.

Description

Return the number of columns present in x.

Usage

h2o.ncol(x)

Arguments

x

An H2OFrame object.

See Also

nrow for the base R implementation, ncol().

Examples

## Not run:
library(h2o)
h2o.init()

iris <- h2o.importFile(f)
h2o.ncol(iris)

## End(Not run)

h2o.networkTest

View Network Traffic Speed

Description

View speed with various file sizes.

Usage

h2o.networkTest()

Value

Returns a table listing the network speed for 1B, 10KB, and 10MB.
h2o.nlevels

*Get the number of factor levels for this frame.*

**Description**

Get the number of factor levels for this frame.

**Usage**

```
h2o.nlevels(x)
```

**Arguments**

- `x`: An H2OFrame object.

**See Also**

`nlevels` for the base R method.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

cars <- h2o.importFile(f)
h2o.nlevels(cars)

## End(Not run)
```

---

h2o.no_progress

*Disable Progress Bar*

**Description**

Disable Progress Bar

**Usage**

```
h2o.no_progress()
```
Examples

## Not run:
library(h2o)
h2o.init()
h2o.no_progress()

iris <- h2o.importFile(f)
iris["class"] <- as.factor(iris["class"])
predictors <- c("sepal_len", "sepal_wid", "petal_len", "petal_wid")
splits <- h2o.splitFrame(iris, ratios = 0.8, seed = 1234)
train <- splits[[1]]
valid <- splits[[2]]

iris_km <- h2o.kmeans(x = predictors,
                      training_frame = train,
                      validation_frame = valid,
                      k = 10, estimate_k = TRUE,
                      standardize = FALSE, seed = 1234)

## End(Not run)

h2o.nrow

Return the number of rows present in x.

Description

Return the number of rows present in x.

Usage

h2o.nrow(x)

Arguments

x An H2OFrame object.

See Also

nrow for the base R implementation.

Examples

## Not run:
library(h2o)
h2o.init()

cars <- h2o.importFile(f)
h2o.null_deviance

Retrieve the null deviance

Description

If "train", "valid", and "xval" parameters are FALSE (default), then the training null deviance value is returned. If more than one parameter is set to TRUE, then a named vector of null deviances are returned, where the names are "train", "valid" or "xval".

Usage

h2o.null_deviance(object, train = FALSE, valid = FALSE, xval = FALSE)

Arguments

object      An H2OModel or H2OModelMetrics
train       Retrieve the training null deviance
valid       Retrieve the validation null deviance
xval        Retrieve the cross-validation null deviance

Examples

## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.importFile(prostate_path)
prostate[, 2] <- as.factor(prostate[, 2])
prostate_glm <- h2o.glm(y = "CAPSULE", x = c("AGE", "RACE", "PSA", "DCAPS"),
                        training_frame = prostate, family = "binomial", nfolds = 0,
                        alpha = 0.5, lambda_search = FALSE)
h2o.null_deviance(prostate_glm, train = TRUE)

## End(Not run)
**h2o.null_dof**

Retrieve the null degrees of freedom.

**Description**

If "train", "valid", and "xval" parameters are FALSE (default), then the training null degrees of freedom value is returned. If more than one parameter is set to TRUE, then a named vector of null degrees of freedom are returned, where the names are "train", "valid" or "xval".

**Usage**

h2o.null_dof(object, train = FALSE, valid = FALSE, xval = FALSE)

**Arguments**

- **object**: An H2OModel or H2OModelMetrics
- **train**: Retrieve the training null degrees of freedom
- **valid**: Retrieve the validation null degrees of freedom
- **xval**: Retrieve the cross-validation null degrees of freedom

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.importFile(prostate_path)
prostate[, 2] <- as.factor(prostate[, 2])
prostate_glm <- h2o.glm(y = "CAPSULE", x = c("AGE", "RACE", "PSA", "DCAPS"),
                        training_frame = prostate, family = "binomial", nfolds = 0,
                        alpha = 0.5, lambda_search = FALSE)
h2o.null_dof(prostate_glm, train = TRUE)
## End(Not run)
```

**h2o.num_iterations**

Retrieve the number of iterations.

**Description**

Retrieve the number of iterations.

**Usage**

h2o.num_iterations(object)
**Arguments**

- **object**: An `H2OClusteringModel` object.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.importFile(prostate_path)
prostate[, 2] <- as.factor(prostate[, 2])
prostate_glm <- h2o.glm(y = "CAPSULE", x = c("AGE", "RACE", "PSA", "DCAPS"),
  training_frame = prostate, family = "binomial",
  nfolds = 0, alpha = 0.5, lambda_search = FALSE)
h2o.num_iterations(prostate_glm)

## End(Not run)
```

---

**h2o.num_valid_substrings**

*Count of substrings >= 2 chars that are contained in file*

---

**Description**

Find the count of all possible substrings >= 2 chars that are contained in the specified line-separated text file.

**Usage**

```r
h2o.num_valid_substrings(x, path)
```

**Arguments**

- **x**: The column on which to calculate the number of valid substrings.
- **path**: Path to text file containing line-separated strings to be referenced.
h2o.openLog

**Description**

Open existing logs of H2O R POST commands and error responses on local disk. Used primarily for debugging purposes.

**Usage**

```r
h2o.openLog(type)
```

**Arguments**

- `type` Currently unimplemented.

**See Also**

h2o.startLogging, h2o.stopLogging, h2o.clearLog

**Examples**

```r
## Not run:
h2o.init()

h2o.startLogging()
australia_path = system.file("extdata", "australia.csv", package = "h2o")
australia = h2o.importFile(path = australia_path)
h2o.stopLogging()

# Not run to avoid windows being opened during R CMD check
# h2o.openLog("Command")
# h2o.openLog("Error")

## End(Not run)
```

h2o.parseRaw

**Description**

The second phase in the data ingestion step.


Usage

```
h2o.parseRaw(
    data,
    pattern = "",  # (Optional) Character string containing a regular expression to match file(s) in the folder.
    destination_frame = "",  # (Optional) The hex key assigned to the parsed file.
    header = NA,  # (Optional) A logical value indicating whether the first row is the column header. If missing, H2O will automatically try to detect the presence of a header.
    sep = "",  # (Optional) The field separator character. Values on each line of the file are separated by this character. If sep = "", the parser will automatically detect the separator.
    col.names = NULL,  # (Optional) An H2OFrame object containing a single delimited line with the column names for the file. If skipped_columns are specified, only list column names of columns that are not skipped.
    col.types = NULL,  # (Optional) A vector specifying the types to attempt to force over columns. If skipped_columns are specified, only list column types of columns that are not skipped.
    na.strings = NULL,  # (Optional) H2O will interpret these strings as missing.
    blocking = FALSE,  # (Optional) Tell H2O parse call to block synchronously instead of polling. This can be faster for small datasets but loses the progress bar.
    parse_type = NULL,  # (Optional) Specify which parser type H2O will use. Valid types are "ARFF", "XLS", "CSV", "SVMLight"
    chunk_size = NULL,  # size of chunk of (input) data in bytes
)
```

Arguments

- **data**: An H2OFrame object to be parsed.
- **pattern**: (Optional) Character string containing a regular expression to match file(s) in the folder.
- **destination_frame**: (Optional) The hex key assigned to the parsed file.
- **header**: (Optional) A logical value indicating whether the first row is the column header. If missing, H2O will automatically try to detect the presence of a header.
- **sep**: (Optional) The field separator character. Values on each line of the file are separated by this character. If sep = "", the parser will automatically detect the separator.
- **col.names**: (Optional) An H2OFrame object containing a single delimited line with the column names for the file. If skipped_columns are specified, only list column names of columns that are not skipped.
- **col.types**: (Optional) A vector specifying the types to attempt to force over columns. If skipped_columns are specified, only list column types of columns that are not skipped.
- **na.strings**: (Optional) H2O will interpret these strings as missing.
- **blocking**: (Optional) Tell H2O parse call to block synchronously instead of polling. This can be faster for small datasets but loses the progress bar.
- **parse_type**: (Optional) Specify which parser type H2O will use. Valid types are "ARFF", "XLS", "CSV", "SVMLight"
h2o.parseSetup

**decrypt_tool**
(Optional) Specify a Decryption Tool (key-reference acquired by calling h2o.decryptionSetup.

**skipped_columns**
a list of column indices to be excluded from parsing

**custom_non_data_line_markers**
(Optional) If a line in imported file starts with any character in given string it will NOT be imported. Empty string means all lines are imported, NULL means that default behaviour for given format will be used

**partition_by**
(Optional) Names of the columns the persisted dataset has been partitioned by.

**quotechar**
A hint for the parser which character to expect as quoting character. None (default) means autodetection.

**escapechar**
(Optional) One ASCII character used to escape other characters.

**Details**

Parse the Raw Data produced by the import phase.

**See Also**

h2o.importFile, h2o.parseSetup

---

**Usage**

```r
h2o.parseSetup(
  data,
  pattern = "",
  destination_frame = "",
  header = NA,
  sep = "",
  col.names = NULL,
  col.types = NULL,
  na.strings = NULL,
  parse_type = NULL,
  chunk_size = NULL,
  decrypt_tool = NULL,
  skipped_columns = NULL,
  custom_non_data_line_markers = NULL,
  partition_by = NULL,
  single_quotes = FALSE,
  escapechar = ""
)
```

---

**Description**

Get a parse setup back for the staged data.
Arguments

- **data**: An H2OFrame object to be parsed.
- **pattern** *(Optional)*: Character string containing a regular expression to match file(s) in the folder.
- **destination_frame** *(Optional)*: The hex key assigned to the parsed file.
- **header** *(Optional)*: A logical value indicating whether the first row is the column header. If missing, H2O will automatically try to detect the presence of a header.
- **sep** *(Optional)*: The field separator character. Values on each line of the file are separated by this character. If sep = "", the parser will automatically detect the separator.
- **col.names** *(Optional)*: An H2OFrame object containing a single delimited line with the column names for the file. If skipped_columns are specified, only list column names of columns that are not skipped.
- **col.types** *(Optional)*: A vector specifying the types to attempt to force over columns. If skipped_columns are specified, only list column types of columns that are not skipped.
- **na.strings** *(Optional)*: H2O will interpret these strings as missing.
- **parse_type** *(Optional)*: Specify which parser type H2O will use. Valid types are "ARFF", "XLS", "CSV", "SVMLight".
- **chunk_size**: Size of chunk of (input) data in bytes.
- **decrypt_tool** *(Optional)*: Specify a Decryption Tool (key-reference acquired by calling h2o.decryptionSetup).
- **skipped_columns**: A list of column indices to be excluded from parsing.
- **custom_non_data_line_markers** *(Optional)*: If a line in imported file starts with any character in given string it will NOT be imported. Empty string means all lines are imported, NULL means that default behaviour for given format will be used.
- **partition_by** *(Optional)*: Names of the columns the persisted dataset has been partitioned by.
- **single_quotes** If set to true, the parser expects single quotes. False for double quotes (default).
- **escapechar** *(Optional)*: One ASCII character used to escape other characters.

See Also

- h2o.parseRaw
**h2o.partialPlot**

**Partial Dependence Plots**

**Description**

Partial dependence plot gives a graphical depiction of the marginal effect of a variable on the response. The effect of a variable is measured in change in the mean response. Note: Unlike randomForest’s partialPlot when plotting partial dependence the mean response (probabilities) is returned rather than the mean of the log class probability.

**Usage**

```r
h2o.partialPlot(
  object,  # An H2OModel object.
  data,  # An H2OFrame object used for scoring and constructing the plot.
  cols,  # Feature(s) for which partial dependence will be calculated.
  destination_key,  # An key reference to the created partial dependence tables in H2O.
  nbins = 20,  # Number of bins used. For categorical columns make sure the number of bins exceeds the level count. If you enable add_missing_NA, the returned length will be nbins+1.
  plot = TRUE,  # A logical specifying whether to plot partial dependence table.
  plot_stddev = TRUE,  # A logical specifying whether to add std err to partial dependence plot.
  weight_column = -1,  # A string denoting which column of data should be used as the weight column.
  include_na = FALSE,  # A logical specifying whether missing value should be included in the Feature values.
  user_splits = NULL,  # A logical specifying whether to add std err to partial dependence plot.
  col_pairs_2dpdp = NULL,  # A logical specifying whether missing value should be included in the Feature values.
  save_to = NULL,  # A logical specifying whether missing value should be included in the Feature values.
  row_index = -1,  # A logical specifying whether missing value should be included in the Feature values.
  targets = NULL  # A logical specifying whether missing value should be included in the Feature values.
)
```

**Arguments**

- **object**: An H2OModel object.
- **data**: An H2OFrame object used for scoring and constructing the plot.
- **cols**: Feature(s) for which partial dependence will be calculated.
- **destination_key**: An key reference to the created partial dependence tables in H2O.
- **nbins**: Number of bins used. For categorical columns make sure the number of bins exceeds the level count. If you enable add_missing_NA, the returned length will be nbins+1.
- **plot**: A logical specifying whether to plot partial dependence table.
- **plot_stddev**: A logical specifying whether to add std err to partial dependence plot.
- **weight_column**: A string denoting which column of data should be used as the weight column.
- **include_na**: A logical specifying whether missing value should be included in the Feature values.
user_splits
A two-level nested list containing user defined split points for pdp plots for each
column. If there are two columns using user defined split points, there should be
two lists in the nested list. Inside each list, the first element is the column name
followed by values defined by the user.

col_pairs_2dpdp
A two-level nested list like this: col_pairs_2dpdp = list(c("col1_name", "col2_name"),
c("col1_name", "col3_name"), ...), where a 2D partial plots will be generated for
col1_name, col2_name pair, for col1_name, col3_name pair and whatever other
pairs that are specified in the nested list.

save_to
Fully qualified prefix of the image files the resulting plots should be saved to,
e.g. '~/home/user/pdp'. Plots for each feature are saved separately in PNG for-
mat, each file receives a suffix equal to the corresponding feature name, e.g.
~/home/user/pdp_AGE.png'. If the files already exists, they will be overridden.
Files are only saves if plot = TRUE (default).

row_index
Row for which partial dependence will be calculated instead of the whole input
frame.

targets
Target classes for multinomial model.

Value
Plot and list of calculated mean response tables for each feature requested.

Examples
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
prostate[, "CAPSULE"] <- as.factor(prostate[, "CAPSULE"] )
prostate[, "RACE"] <- as.factor(prostate[, "RACE"] )
prostate_gbm <- h2o.gbm(x = c("AGE", "RACE"),
y = "CAPSULE",
  training_frame = prostate,
  ntrees = 10,
  max_depth = 5,
  learn_rate = 0.1)
h2o.partialPlot(object = prostate_gbm, data = prostate, cols = c("AGE", "RACE"))

iris_hex <- as.h2o(iris)
iris_gbm <- h2o.gbm(x = c(1:4), y = 5, training_frame = iris_hex)

# one target class
h2o.partialPlot(object = iris_gbm, data = iris_hex, cols="Petal.Length", targets=c("setosa"))
# three target classes
h2o.partialPlot(object = iris_gbm, data = iris_hex, cols="Petal.Length",
  targets=c("setosa", "virginica", "versicolor"))

## End(Not run)
**h2o.pd_multi_plot**

*Plot partial dependencies for a variable across multiple models*

**Description**
Partial dependence plot (PDP) gives a graphical depiction of the marginal effect of a variable on the response. The effect of a variable is measured in change in the mean response. PDP assumes independence between the feature for which is the PDP computed and the rest.

**Usage**

```r
h2o.pd_multi_plot(
  object,
  newdata,
  column,
  best_of_family = TRUE,
  target = NULL,
  row_index = NULL,
  max_levels = 30
)
```

**Arguments**
- `object` Either a list of H2O models/model_ids or an H2OAutoML object.
- `newdata` An H2OFrame.
- `column` A feature column name to inspect. Character string.
- `best_of_family` If TRUE, plot only the best model of each algorithm family; if FALSE, plot all models. Defaults to TRUE.
- `target` If multinomial, plot PDP just for `target` category.
- `row_index` Optional. Calculate Individual Conditional Expectation (ICE) for row, `row_index`. Integer.
- `max_levels` An integer specifying the maximum number of factor levels to show. Defaults to 30.

**Value**
A ggplot2 object

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

# Import the wine dataset into H2O:
```
df <- h2o.importFile(f)

# Set the response
response <- "quality"

# Split the dataset into a train and test set:
splits <- h2o.splitFrame(df, ratios = 0.8, seed = 1)
train <- splits[[1]]
test <- splits[[2]]

# Build and train the model:
aml <- h2o.automl(y = response,
                 training_frame = train,
                 max_models = 10,
                 seed = 1)

# Create the partial dependence plot
pdp <- h2o.pd_multi_plot(aml, test, column = "alcohol")
print(pdp)

## End(Not run)

---

**h2o.pd_plot**

*Plot partial dependence for a variable*

**Description**

Partial dependence plot (PDP) gives a graphical depiction of the marginal effect of a variable on the response. The effect of a variable is measured in change in the mean response. PDP assumes independence between the feature for which is the PDP computed and the rest.

**Usage**

```r
h2o.pd_plot(
  object,  
  newdata,  
  column,  
  target = NULL,  
  row_index = NULL,  
  max_levels = 30
)
```

**Arguments**

- `object` An H2O model.
- `newdata` An H2OFrame. Used to generate predictions used in Partial Dependence calculations.
- `column` A feature column name to inspect. Character string.
### target
If multinomial, plot PDP just for target category. Character string.

### row_index
Optional. Calculate Individual Conditional Expectation (ICE) for row, row_index. Integer.

### max_levels
An integer specifying the maximum number of factor levels to show. Defaults to 30.

#### Value
A ggplot2 object

#### Examples
```r
## Not run:
library(h2o)
h2o.init()

# Import the wine dataset into H2O:
df <- h2o.importFile(f)

# Set the response
target <- "quality"

# Split the dataset into a train and test set:
splits <- h2o.splitFrame(df, ratios = 0.8, seed = 1)
train <- splits[[1]]
test <- splits[[2]]

# Build and train the model:
gbm <- h2o.gbm(y = target, training_frame = train)

# Create the partial dependence plot
pdp <- h2o.pd_plot(gbm, test, column = "alcohol")
print(pdp)
```

## Model Performance Metrics in H2O

### Description
Given a trained h2o model, compute its performance on the given dataset. However, if the dataset
does not contain the response/target column, no performance will be returned. Instead, a warning
message will be printed.
Usage

```r
h2o.performance(
  model,
  newdata = NULL,
  train = FALSE,
  valid = FALSE,
  xval = FALSE,
  data = NULL,
  auc_type = "NONE"
)
```

Arguments

- **model**: An H2OModel object
- **newdata**: An H2OFrame. The model will make predictions on this dataset, and subsequently score them. The dataset should match the dataset that was used to train the model, in terms of column names, types, and dimensions. If newdata is passed in, then train, valid, and xval are ignored.
- **train**: A logical value indicating whether to return the training metrics (constructed during training). Note: when the trained h2o model uses balance_classes, the training metrics constructed during training will be from the balanced training dataset. For more information visit: [https://0xdata.atlassian.net/browse/TN-9](https://0xdata.atlassian.net/browse/TN-9)
- **valid**: A logical value indicating whether to return the validation metrics (constructed during training).
- **xval**: A logical value indicating whether to return the cross-validation metrics (constructed during training).
- **data** (DEPRECATED): An H2OFrame. This argument is now called `newdata`.
- **auc_type**: For multinomila model only. Set default multinomial AUC type. Must be one of: "AUTO", "NONE", "MACRO_OVR", "WEIGHTED_OVR", "MACRO_OVO", "WEIGHTED_OVO". Default is "NONE"

Value

Returns an object of the H2OModelMetrics subclass.

Examples

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
prostate$CAPSULE <- as.factor(prostate$CAPSULE)
prostate_gbm <- h2o.gbm(3:9, "CAPSULE", prostate)
h2o.performance(model = prostate_gbm, newdata=prostate)
```
## If model uses balance_classes
## the results from train = TRUE will not match the results from newdata = prostate
prostate_gbm_balanced <- h2o.gbm(3:9, "CAPSULE", prostate, balance_classes = TRUE)
h2o.performance(model = prostate_gbm_balanced, newdata = prostate)
h2o.performance(model = prostate_gbm_balanced, train = TRUE)

## End(Not run)

---

**h2o.permutation_importance**

*Calculate Permutation Feature Importance.*

### Description

When `n_repeats == 1`, the result is similar to the one from `h2o.varimp()`, i.e., it contains the following columns "Relative Importance", "Scaled Importance", and "Percentage".

### Usage

```r
h2o.permutation_importance(
  object,
  newdata,
  metric = c("AUTO", "AUC", "MAE", "MSE", "RMSE", "logloss", "mean_per_class_error", "PR_AUC"),
  n_samples = 10000,
  n_repeats = 1,
  features = NULL,
  seed = -1
)
```

### Arguments

- **object**: A trained supervised H2O model.
- **newdata**: Training frame of the model which is going to be permuted.
- **metric**: Metric to be used. One of "AUTO", "AUC", "MAE", "MSE", "RMSE", "logloss", "mean_per_class_error", "PR_AUC". Defaults to "AUTO".
- **n_samples**: Number of samples to be evaluated. Use -1 to use the whole dataset. Defaults to 10 000.
- **n_repeats**: Number of repeated evaluations. Defaults to 1.
- **features**: Character vector of features to include in the permutation importance. Use NULL to include all.
- **seed**: Seed for the random generator. Use -1 to pick a random seed. Defaults to -1.
Details

When \( n_{repeats} > 1 \), the individual columns correspond to the permutation variable importance values from individual runs which corresponds to the "Relative Importance" and also to the distance between the original prediction error and prediction error using a frame with a given feature permuted.

Value

H2OTable with variable importance.

Examples

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.importFile(prostate_path)
prostate[, 2] <- as.factor(prostate[, 2])
model <- h2o.gbm(x = 3:9, y = 2, training_frame = prostate, distribution = "bernoulli")
h2o.permutation_importance(model, prostate)
## End(Not run)
```

h2o.permutation_importance_plot

**Plot Permutation Variable Importances.**

Description

This method plots either a bar plot or if \( n_{repeats} > 1 \) a box plot and returns the variable importance table.

Usage

```r
h2o.permutation_importance_plot(
  object,
  newdata,
  metric = c("AUTO", "AUC", "MAE", "MSE", "RMSE", "logloss", "mean_per_class_error", "PR_AUC"),
  n_samples = 10000,
  n_repeats = 1,
  features = NULL,
  seed = -1,
  num_of_features = NULL
)
```
h2o.pivot

Pivot a frame

Description

Pivot the frame designated by the three columns: index, column, and value. Index and column should be of type enum, int, or time. For cases of multiple indexes for a column label, the aggregation method is to pick the first occurrence in the data frame.

Usage

h2o.pivot(x, index, column, value)
Arguments

x  an H2OFrame
index  the column where pivoted rows should be aligned on
column  the column to pivot
value  values of the pivoted table

Value

An H2OFrame with columns from the columns arg, aligned on the index arg, with values from values arg

Examples

## Not run:
library(h2o)
h2o.init()

df = h2o.createFrame(rows = 1000, cols = 3, factors = 10, integer_fraction = 1.0/3,
categorical_fraction = 1.0/3, missing_fraction = 0.0, seed = 123)
df$C3 = h2o.abs(df$C3)
h2o.pivot(df, index="C3", column="C2", value="C1")

## End(Not run)

h2o.prcomp

Principal component analysis of an H2O data frame

Description

Principal components analysis of an H2O data frame using the power method to calculate the singular value decomposition of the Gram matrix.

Usage

h2o.prcomp(
  training_frame,
  x,
  model_id = NULL,
  validation_frame = NULL,
  ignore_const_cols = TRUE,
  score_each_iteration = FALSE,
  transform = c("NONE", "STANDARDIZE", "NORMALIZE", "DEMEAN", "DESCALE"),
  pca_method = c("GramSVD", "Power", "Randomized", "GLRM"),
  pca_impl = c("MTJ_EVD_DENSEMATRIX", "MTJ_EVD_SYMMATRIX", "MTJ_SVD_DENSEMATRIX",
              "JAMA"),
  k = 1,
  max_iterations = 1000,
Arguments

**training_frame**  
Id of the training data frame.

**x**  
A vector containing the character names of the predictors in the model.

**model_id**  
Destination id for this model; auto-generated if not specified.

**validation_frame**  
Id of the validation data frame.

**ignore_const_cols**  
Logical. Ignore constant columns. Defaults to TRUE.

**score_each_iteration**  
Logical. Whether to score during each iteration of model training. Defaults to FALSE.

**transform**  
Transformation of training data Must be one of: "NONE", "STANDARDIZE", "NORMALIZE", "DEMEAN", "DESCALE". Defaults to NONE.

**pca_method**  
Specify the algorithm to use for computing the principal components: GramSVD - uses a distributed computation of the Gram matrix, followed by a local SVD; Power - computes the SVD using the power iteration method (experimental); Randomized - uses randomized subspace iteration method; GLRM - fits a generalized low-rank model with L2 loss function and no regularization and solves for the SVD using local matrix algebra (experimental) Must be one of: "GramSVD", "Power", "Randomized", "GLRM". Defaults to GramSVD.

**pca_impl**  

**k**  
Rank of matrix approximation Defaults to 1.

**max_iterations**  
Maximum training iterations Defaults to 1000.

**use_all_factor_levels**  
Logical. Whether first factor level is included in each categorical expansion Defaults to FALSE.

**compute_metrics**  
Logical. Whether to compute metrics on the training data Defaults to TRUE.

**impute_missing**  
Logical. Whether to impute missing entries with the column mean Defaults to FALSE.
h2o.predict

Predict on an H2O Model

Description

Predict on an H2O Model

Usage

h2o.predict(object, newdata, ...)

seed

Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).

max_runtime_secs

Maximum allowed runtime in seconds for model training. Use 0 to disable. Defaults to 0.

export_checkpoints_dir

Automatically export generated models to this directory.

Value

an object of class H2ODimReductionModel.

References


See Also

h2o.svd, h2o.glrm

Examples

## Not run:
library(h2o)
h2o.init()
australia_path <- system.file("extdata", "australia.csv", package = "h2o")
australia <- h2o.uploadFile(path = australia_path)
h2o.prcomp(training_frame = australia, k = 8, transform = "STANDARDIZE")

## End(Not run)
**Arguments**

- **object**: a fitted model object for which prediction is desired.
- **newdata**: An H2OFrame object in which to look for variables with which to predict.
- **...**: additional arguments to pass on.

**Value**

Returns an H2OFrame object with probabilites and default predictions.

---

**Description**

Provides the method `h2o.predict` with which you can predict a MOJO or POJO Jar model from R.

**Usage**

```r
h2o.predict_json(model, json, genmodelpath, labels, classpath, javaoptions)
```

**Arguments**

- **model**: String with file name of MOJO or POJO Jar
- **json**: JSON String with inputs to model
- **genmodelpath**: (Optional) path name to h2o-genmodel.jar, if not set defaults to same dir as MOJO
- **labels**: (Optional) if TRUE then show output labels in result
- **classpath**: (Optional) Extra items for the class path of where to look for Java classes, e.g., h2o-genmodel.jar
- **javaoptions**: (Optional) Java options string, default if "-Xmx4g"

**Value**

Returns an object with the prediction result

**Examples**

```r
## Not run:
library(h2o)
h2o.predict_json(~/GBM_model_python_1473313897851_6.zip, '{"C7":1}')
h2o.predict_json(~/GBM_model_python_1473313897851_6.zip, '{"C7":1}', c('.', 'lib'))

## End(Not run)
```
h2o.predict_rules

Evaluates validity of the given rules on the given data. Returns a frame with a column per each input rule id, representing a flag whether given rule is applied to the observation or not.

Description

Evaluates validity of the given rules on the given data. Returns a frame with a column per each input rule id, representing a flag whether given rule is applied to the observation or not.

Usage

h2o.predict_rules(model, frame, rule_ids)

Arguments

model A trained rulefit model.
frame A frame on which rule validity is to be evaluated
rule_ids Rule ids to be evaluated against the frame

Examples

```r
## Not run:
library(h2o)
h2o.init()
titanic <- h2o.importFile(
  "https://s3.amazonaws.com/h2o-public-test-data/smalldata/gbm_test/titanic.csv"
)
response = "survived"
predictors <- c("age", "sibsp", "parch", "fare", "sex", "pclass")
titanic[,response] <- as.factor(titanic[,response])
titanic[,"pclass"] <- as.factor(titanic[,"pclass"])
splits <- h2o.splitFrame(data = titanic, ratios = .8, seed = 1234)
train <- splits[[1]]
test <- splits[[2]]

rfit <- h2o.rulefit(y = response, x = predictors, training_frame = train, validation_frame = test, 
  min_rule_length = 1, max_rule_length = 10, max_num_rules = 100, seed = 1, model_type="rules")
h2o.predict_rules(rfit, train, c("M1T0N7", "M1T49N7", "M1T16N7", "M1T36N7", "M2T19N19"))

## End(Not run)
```
h2o.print

Print An H2OFrame

Description

Print An H2OFrame

Usage

h2o.print(x, n = 6L)

Arguments

x An H2OFrame object
n An (Optional) A single integer. If positive, number of rows in x to return. If negative, all but the n first/last number of rows in x. Anything bigger than 20 rows will require asking the server (first 20 rows are cached on the client).

Examples

```r
## Not run:
library()
h2o.init()

iris <- h2o.importFile(f)
h2o.print(iris["species"], n = 15)
## End(Not run)
```

h2o.prod

Return the product of all the values present in its arguments.

Description

Return the product of all the values present in its arguments.

Usage

h2o.prod(x)

Arguments

x An H2OFrame object.
See Also

prod for the base R implementation.

Examples

```r
## Not run:
library(h2o)
h2o.init()

iris <- h2o.importFile(f)
h2o.prod(iris["petal_len"])

## End(Not run)
```

---

**h2o.proj_archetypes**  
Convert Archetypes to Features from H2O GLRM Model

### Description

Project each archetype in an H2O GLRM model into the corresponding feature space from the H2O training frame.

### Usage

```
h2o.proj_archetypes(object, data, reverse_transform = FALSE)
```

### Arguments

- **object**: An `H2ODimReductionModel` object that represents the model containing archetypes to be projected.
- **data**: An `H2OFrame` object representing the training data for the H2O GLRM model.
- **reverse_transform**: (Optional) A logical value indicating whether to reverse the transformation from model-building by re-scaling columns and adding back the offset to each column of the projected archetypes.

### Value

Returns an `H2OFrame` object containing the projection of the archetypes down into the original feature space, where each row is one archetype.

### See Also

h2o.glrm for making an `H2ODimReductionModel`.  

## Examples

```r
## Not run:
library(h2o)
h2o.init()
iris_hf <- as.h2o(iris)
iris_glrm <- h2o.glrm(training_frame = iris_hf, k = 4, loss = "Quadratic",
                      multi_loss = "Categorical", max_iterations = 1000)
iris_parch <- h2o.proj_archetypes(iris_glrm, iris_hf)
head(iris_parch)
## End(Not run)
```

---

**h2o.psvm**  
*Trains a Support Vector Machine model on an H2O dataset*

---

**Description**

Alpha version. Supports only binomial classification problems.

**Usage**

```r
h2o.psvm(
  x,
  y,
  training_frame,
  model_id = NULL,
  validation_frame = NULL,
  ignore_const_cols = TRUE,
  hyper_param = 1,
  kernel_type = c("gaussian"),
  gamma = -1,
  rank_ratio = -1,
  positive_weight = 1,
  negative_weight = 1,
  disable_training_metrics = TRUE,
  sv_threshold = 1e-04,
  fact_threshold = 1e-05,
  feasible_threshold = 0.001,
  surrogate_gap_threshold = 0.001,
  mu_factor = 10,
  max_iterations = 200,
  seed = -1
)
```

**Arguments**

- `x`  
  (Optional) A vector containing the names or indices of the predictor variables to use in building the model. If `x` is missing, then all columns except `y` are used.
The name or column index of the response variable in the data. The response must be either a binary categorical/factor variable or a numeric variable with values -1/1 (for compatibility with SVMlight format).

Id of the training data frame.

Destination id for this model; auto-generated if not specified.

Id of the validation data frame.

Logical. Ignore constant columns. Defaults to TRUE.

Penalty parameter C of the error term Defaults to 1.

Type of used kernel Must be one of: "gaussian". Defaults to gaussian.

Coefficient of the kernel (currently RBF gamma for gaussian kernel, -1 means 1/#features) Defaults to -1.

Desired rank of the ICF matrix expressed as a ration of number of input rows (-1 means use sqrt(#rows)). Defaults to -1.

Weight of positive (+1) class of observations Defaults to 1.

Weight of positive (-1) class of observations Defaults to 1.

Logical. Disable calculating training metrics (expensive on large datasets) Defaults to TRUE.

Threshold for accepting a candidate observation into the set of support vectors Defaults to 0.0001.

Convergence threshold of the Incomplete Cholesky Factorization (ICF) Defaults to 1e-05.

Convergence threshold for primal-dual residuals in the IPM iteration Defaults to 0.001.

Feasibility criterion of the surrogate duality gap (eta) Defaults to 0.001.

Increasing factor mu Defaults to 10.

Maximum number of iteration of the algorithm Defaults to 200.

Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).

Examples

```r
## Not run:
library(h2o)
h2o.init()

# Import the splice dataset
f <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/splice/splice.svm"
```
splice <- h2o.importFile(f)

# Train the Support Vector Machine model
svm_model <- h2o.psvm(gamma = 0.01, rank_ratio = 0.1,
   y = "C1", training_frame = splice,
   disable_training_metrics = FALSE)

## End(Not run)

---

**h2o.quantile**

**Quantiles of H2O Frames.**

**Description**

Obtain and display quantiles for H2O parsed data.

**Usage**

```r
h2o.quantile(
  x,
  probs = c(0.001, 0.01, 0.1, 0.25, 0.333, 0.5, 0.667, 0.75, 0.9, 0.99, 0.999),
  combine_method = c("interpolate", "average", "avg", "low", "high"),
  weights_column = NULL,
  ...
)

## S3 method for class 'H2OFrame'
quantile(
  x,
  probs = c(0.001, 0.01, 0.1, 0.25, 0.333, 0.5, 0.667, 0.75, 0.9, 0.99, 0.999),
  combine_method = c("interpolate", "average", "avg", "low", "high"),
  weights_column = NULL,
  ...
)
```

**Arguments**

- `x`: An H2OFrame object with a single numeric column.
- `probs`: Numeric vector of probabilities with values in [0,1].
- `combine_method`: How to combine quantiles for even sample sizes. Default is to do linear interpolation. E.g., If method is "lo", then it will take the lo value of the quantile. Abbreviations for average, low, and high are acceptable (avg, lo, hi).
- `weights_column`: (Optional) String name of the observation weights column in x or an H2OFrame object with a single numeric column of observation weights.
- `...`: Further arguments passed to or from other methods.
Details


Value

A vector describing the percentiles at the given cutoffs for the H2OFrame object.

Examples

```r
## Not run:
# Request quantiles for an H2O parsed data set:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
# Request quantiles for a subset of columns in an H2O parsed data set
quantile(prostate[, 3])
for(i in 1:ncol(prostate))
  quantile(prostate[, i])
## End(Not run)
```

h2o.r2

Retrieves the R2 value from an H2O model. Will return R^2 for GLM Models and will return NaN otherwise. If "train", "valid", and "xval" parameters are FALSE (default), then the training R2 value is returned. If more than one parameter is set to TRUE, then a named vector of R2s are returned, where the names are "train", "valid" or "xval".

Usage

`h2o.r2(object, train = FALSE, valid = FALSE, xval = FALSE)`

Arguments

- `object` An `H2OModel` object.
- `train` Retrieve the training R2
- `valid` Retrieve the validation set R2 if a validation set was passed in during model build time.
- `xval` Retrieve the cross-validation R2
### Examples

```r
## Not run:
library(h2o)

h <- h2o.init()
fr <- as.h2o(iris)

m <- h2o.glm(x = 2:5, y = 1, training_frame = fr)

h2o.r2(m)

## End(Not run)
```

---

### h2o.randomForest

Build a Random Forest model

### Description

Builds a Random Forest model on an H2OFrame.

### Usage

```r
h2o.randomForest(
  x,
  y,
  training_frame,
  model_id = NULL,
  validation_frame = NULL,
  nfolds = 0,
  keep_cross_validation_models = TRUE,
  keep_cross_validation_predictions = FALSE,
  keep_cross_validation_fold_assignment = FALSE,
  score_each_iteration = FALSE,
  score_tree_interval = 0,
  fold_assignment = c("AUTO", "Random", "Modulo", "Stratified"),
  fold_column = NULL,
  ignore_const_cols = TRUE,
  offset_column = NULL,
  weights_column = NULL,
  balance_classes = FALSE,
  class_sampling_factors = NULL,
  max_after_balance_size = 5,
  ntrees = 50,
  max_depth = 20,
  min_rows = 1,
  nbins = 20,
  nbins_top_level = 1024,
)```
nbins_cats = 1024,
r2_stopping = Inf,
stopping_rounds = 0,
stopping_metric = c("AUTO", "deviance", "logloss", "MSE", "RMSE", "MAE", "RMSLE", "AUC", "AUCPR", "lift_top_group", "misclassification", "mean_per_class_error", "custom", "custom_increasing"),
stopping_tolerance = 0.001,
max_runtime_secs = 0,
seed = -1,
build_tree_one_node = FALSE,
mtries = -1,
sample_rate = 0.632,
sample_rate_per_class = NULL,
binomial_double_trees = FALSE,
checkpoint = NULL,
col_sample_rate_change_per_level = 1,
col_sample_rate_change_per_tree = 1,
min_split_improvement = 1e-05,
histogram_type = c("AUTO", "UniformAdaptive", "Random", "QuantilesGlobal", "RoundRobin"),
calibrate_model = FALSE,
calibration_frame = NULL,
distribution = c("AUTO", "bernoulli", "multinomial", "gaussian", "poisson", "gamma", "tweedie", "laplace", "quantile", "huber"),
custom_metric_func = NULL,
export_checkpoints_dir = NULL,
check_constant_response = TRUE,
gainslift_bins = -1,
auc_type = c("AUTO", "NONE", "MACRO_OVR", "WEIGHTED_OVR", "MACRO_OVO", "WEIGHTED_OVO"),
verbose = FALSE}

Arguments

x (Optional) A vector containing the names or indices of the predictor variables to use in building the model. If x is missing, then all columns except y are used.

y The name or column index of the response variable in the data. The response must be either a numeric or a categorical/factor variable. If the response is numeric, then a regression model will be trained, otherwise it will train a classification model.

training_frame Id of the training data frame.
model_id Destination id for this model; auto-generated if not specified.
validation_frame Id of the validation data frame.
n folds | Number of folds for K-fold cross-validation (0 to disable or >= 2). Defaults to 0.
keep_cross_validation_models | Logical. Whether to keep the cross-validation models. Defaults to TRUE.
keep_cross_validation_predictions | Logical. Whether to keep the predictions of the cross-validation models. Defaults to FALSE.
keep_cross_validation_fold_assignment | Logical. Whether to keep the cross-validation fold assignment. Defaults to FALSE.
score_each_iteration | Logical. Whether to score during each iteration of model training. Defaults to FALSE.
score_tree_interval | Score the model after every so many trees. Disabled if set to 0. Defaults to 0.
fold_assignment | Cross-validation fold assignment scheme, if fold_column is not specified. The 'Stratified' option will stratify the folds based on the response variable, for classification problems. Must be one of: "AUTO", "Random", "Modulo", "Stratified". Defaults to AUTO.
fold_column | Column with cross-validation fold index assignment per observation.
ignore_const_cols | Logical. Ignore constant columns. Defaults to TRUE.
offset_column | Offset column. This argument is deprecated and has no use for Random Forest.
weights_column | Column with observation weights. Giving some observation a weight of zero is equivalent to excluding it from the dataset; giving an observation a relative weight of 2 is equivalent to repeating that row twice. Negative weights are not allowed. Note: Weights are per-row observation weights and do not increase the size of the data frame. This is typically the number of times a row is repeated, but non-integer values are supported as well. During training, rows with higher weights matter more, due to the larger loss function pre-factor. If you set weight = 0 for a row, the returned prediction frame at that row is zero and this is incorrect. To get an accurate prediction, remove all rows with weight == 0.
balance_classes | Logical. Balance training data class counts via over/under-sampling (for imbalanced data). Defaults to FALSE.
class_sampling_factors | Desired over/under-sampling ratios per class (in lexicographic order). If not specified, sampling factors will be automatically computed to obtain class balance during training. Requires balance_classes.
max_after_balance_size | Maximum relative size of the training data after balancing class counts (can be less than 1.0). Requires balance_classes. Defaults to 5.0.
ntrees | Number of trees. Defaults to 50.
max_depth | Maximum tree depth (0 for unlimited). Defaults to 20.
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>min_rows</td>
<td>Fewest allowed (weighted) observations in a leaf. Defaults to 1.</td>
</tr>
<tr>
<td>nbins</td>
<td>For numerical columns (real/int), build a histogram of (at least) this many bins, then split at the best point. Defaults to 20.</td>
</tr>
<tr>
<td>nbins_top_level</td>
<td>For numerical columns (real/int), build a histogram of (at most) this many bins at the root level, then decrease by factor of two per level. Defaults to 1024.</td>
</tr>
<tr>
<td>nbins_cats</td>
<td>For categorical columns (factors), build a histogram of this many bins, then split at the best point. Higher values can lead to more overfitting. Defaults to 1024.</td>
</tr>
<tr>
<td>r2_stopping</td>
<td>r2_stopping is no longer supported and will be ignored if set - please use stopping_rounds, stopping_metric and stopping_tolerance instead. Previous version of H2O would stop making trees when the ( R^2 ) metric equals or exceeds this. Defaults to ( 1.797693135 \times 10^{308} ).</td>
</tr>
<tr>
<td>stopping_rounds</td>
<td>Early stopping based on convergence of stopping_metric. Stop if simple moving average of length ( k ) of the stopping_metric does not improve for ( k = \text{stopping} _\text{rounds} ). Defaults to 0.</td>
</tr>
<tr>
<td>stopping_metric</td>
<td>Metric to use for early stopping (AUTO: logloss for classification, deviance for regression and anomaly_score for Isolation Forest). Note that custom and custom_increasing can only be used in GBM and DRF with the Python client. Must be one of: &quot;AUTO&quot;, &quot;deviance&quot;, &quot;logloss&quot;, &quot;MSE&quot;, &quot;RMSE&quot;, &quot;MAE&quot;, &quot;RMSLE&quot;, &quot;AUC&quot;, &quot;AUCPR&quot;, &quot;lift_top_group&quot;, &quot;misclassification&quot;, &quot;mean_per_class_error&quot;, &quot;custom&quot;, &quot;custom_increasing&quot;. Defaults to AUTO.</td>
</tr>
<tr>
<td>stopping_tolerance</td>
<td>Relative tolerance for metric-based stopping criterion (stop if relative improvement is not at least this much). Defaults to 0.001.</td>
</tr>
<tr>
<td>max_runtime_secs</td>
<td>Maximum allowed runtime in seconds for model training. Use 0 to disable. Defaults to 0.</td>
</tr>
<tr>
<td>seed</td>
<td>Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).</td>
</tr>
<tr>
<td>build_tree_one_node</td>
<td>Logical. Run on one node only; no network overhead but fewer cpus used. Suitable for small datasets. Defaults to FALSE.</td>
</tr>
<tr>
<td>mtries</td>
<td>Number of variables randomly sampled as candidates at each split. If set to -1, defaults to ( \sqrt{p} ) for classification and ( p/3 ) for regression (where ( p ) is the # of predictors). Defaults to -1.</td>
</tr>
<tr>
<td>sample_rate</td>
<td>Row sample rate per tree (from 0.0 to 1.0). Defaults to 0.632.</td>
</tr>
<tr>
<td>sample_rate_per_class</td>
<td>A list of row sample rates per class (relative fraction for each class, from 0.0 to 1.0), for each tree</td>
</tr>
<tr>
<td>binomial_double_trees</td>
<td>Logical. For binary classification: Build 2x as many trees (one per class) - can lead to higher accuracy. Defaults to FALSE.</td>
</tr>
</tbody>
</table>
h2o.randomForest

checkpoint  Model checkpoint to resume training with.
col_sample_rate_change_per_level
  Relative change of the column sampling rate for every level (must be > 0.0 and <= 2.0). Defaults to 1.
col_sample_rate_per_tree
  Column sample rate per tree (from 0.0 to 1.0). Defaults to 1.
min_split_improvement
  Minimum relative improvement in squared error reduction for a split to happen. Defaults to 1e-05.
histogram_type
  What type of histogram to use for finding optimal split points. Must be one of: "AUTO", "UniformAdaptive", "Random", "QuantilesGlobal", "RoundRobin". Defaults to AUTO.
categorical_encoding
  Encoding scheme for categorical features. Must be one of: "AUTO", "Enum", "OneHotInternal", "OneHotExplicit", "Binary", "Eigen", "LabelEncoder", "Sort-ByResponse", "EnumLimited". Defaults to AUTO.
calibrate_model
  Logical. Use Platt Scaling to calculate calibrated class probabilities. Calibration can provide more accurate estimates of class probabilities. Defaults to FALSE.
calibration_frame
  Calibration frame for Platt Scaling.
distribution
  Distribution. This argument is deprecated and has no use for Random Forest.
custom_metric_func
  Reference to custom evaluation function, format: 'language:keyName=funcName'.
export_checkpoints_dir
  Automatically export generated models to this directory.
check_constant_response
  Logical. Check if response column is constant. If enabled, then an exception is thrown if the response column is a constant value. If disabled, then the model will train regardless of the response column being a constant value or not. Defaults to TRUE.
gainslift_bins
  Gains/Lift table number of bins. 0 means disabled. Default value -1 means automatic binning. Defaults to -1.
auc_type
  Set default multinomial AUC type. Must be one of: "AUTO", "NONE", "MACRO_OVR", "WEIGHTED_OVR", "MACRO_OVO", "WEIGHTED_OVO". Defaults to AUTO.
verbose
  Logical. Print scoring history to the console (Metrics per tree). Defaults to FALSE.

Value

Creates a H2OModel object of the right type.

See Also

predict.H2OModel for prediction
Examples

```r
## Not run:
library(h2o)
h2o.init()

# Import the cars dataset
cars <- h2o.importFile(f)

# Set predictors and response; set response as a factor
cars["economy_20mpg"] <- as.factor(cars["economy_20mpg"])
predictors <- c("displacement", "power", "weight", "acceleration", "year")
response <- "economy_20mpg"

# Train the DRF model
cars_drf <- h2o.randomForest(x = predictors, y = response,
                             training_frame = cars, nfolds = 5,
                             seed = 1234)

## End(Not run)
```

h2o.range

**h2o.range**

Returns a vector containing the minimum and maximum of all the given arguments.

Description

Returns a vector containing the minimum and maximum of all the given arguments.

Usage

h2o.range(x, na.rm = FALSE, finite = FALSE)

Arguments

- **x**: An H2OFrame object.
- **na.rm**: logical. indicating whether missing values should be removed.
- **finite**: logical. indicating if all non-finite elements should be omitted.

See Also

`range` for the base R implementation.
h2o.rank_within_group_by

Examples

```r
## Not run:
library(h2o)
h2o.init()
iris <- h2o.importFile(f)
h2o.range(iris["petal_len"], na.rm = TRUE, finite = TRUE)

## End(Not run)
```

h2o.rank_within_group_by

This function will add a new column rank where the ranking is produced as follows: 1. sorts the H2OFrame by columns specified in group_by_cols and sort_cols in the directions specified by the ascending for the sort_cols. The sort directions for the group_by_cols are ascending only. 2. A new rank column is added to the frame which will contain a rank assignment performed next. The user can choose to assign a name to this new column. The default name is New_Rank_column. 3. For each groupby groups, a rank is assigned to the row starting from 1, 2, ... to the end of that group. 4. If sort_cols_sorted is TRUE, a final sort on the frame will be performed frame according to the sort_cols and the sort directions in ascending. If sort_cols_sorted is FALSE (by default), the frame from step 3 will be returned as is with no extra sort. This may provide a small speedup if desired.

Description

This function will add a new column rank where the ranking is produced as follows: 1. sorts the H2OFrame by columns specified in group_by_cols and sort_cols in the directions specified by the ascending for the sort_cols. The sort directions for the group_by_cols are ascending only. 2. A new rank column is added to the frame which will contain a rank assignment performed next. The user can choose to assign a name to this new column. The default name is New_Rank_column. 3. For each groupby groups, a rank is assigned to the row starting from 1, 2, ... to the end of that group. 4. If sort_cols_sorted is TRUE, a final sort on the frame will be performed frame according to the sort_cols and the sort directions in ascending. If sort_cols_sorted is FALSE (by default), the frame from step 3 will be returned as is with no extra sort. This may provide a small speedup if desired.

Usage

```r
h2o.rank_within_group_by(
  x,
  group_by_cols,
  sort_cols,
)```
h2o.rank_within_group_by

ascending = NULL,
new_col_name = "New_Rank_column",
sort_cols_sorted = FALSE
)

Arguments

x
The H2OFrame input to be sorted.
group_by_cols
a list of column names or indices to form the groupby groups
sort_cols
a list of column names or indices for sorting
ascending
a list of Boolean to determine if ascending sort (set to TRUE) is needed for each
column in sort_cols (optional). Default is ascending sort for all. To perform
descending sort, set value to FALSE
new_col_name
new column name for the newly added rank column if specified (optional). De-
fault name is New_Rank_column.
sort_cols_sorted
Boolean to determine if the final returned frame is to be sorted according to the
sort_cols and sort directions in ascending. Default is FALSE.

The following example is generated by Nidhi Mehta.

If the input frame is train:

ID Group_by_column num data Column_to_arrange_by num_1 fdata 12 1 2941.552 1 3 -3177.9077 1 12 1 2941.552 1 5 -13311.8247 1 12 2 -22722.174 1 3 -3177.9077 1 12 2 -22722.174 1 5 -13311.8247 1 13 3 -12776.884 1 5 -18421.6171 0 13 3 -12776.884 1 4 28080.1607 0 13 1 -6049.830 1 5 -18421.6171 0 13 1 -6049.830 1 4 28080.1607 0 15 3 -16995.346 1 1 -9781.6373 0 16 1 -1003.593 0 3 -61284.6900 0 16 3 -22905.288 0 3 -61284.6900 0 16 3 -22905.288 0 3 -61284.6900 0 17 2 -13465.496 1 2 12094.4851 1 17 2 -13465.496 1 3 -415.1114 0 17 2 -13465.496 1 3 -415.1114 0 17 2 -3329.619 1 3 -11772.1338 1 17 2 -3329.619 1 3 -415.1114 0

If the following commands are issued: rankedF1 <- h2o.rank_within_group_by(train,
c("Group_by_column"), c("Column_to_arrange_by"), c(TRUE))
h2o.summary(rankedF1)
The returned frame rankedF1 will look like this: ID Group_by_column num fdata.1 Column_to_arrange_by num_1 fdata.1 1 3 -3177.9077 1 11 6 1 -1003.593 0 3 -61284.6900 0 12 1 -6049.830 0 4 28080.1607 0 3 12 1 2941.552 1 5 -13311.8247 1 4 13 1 -6049.830 0 5 -18421.6171 0 5 17 2 -13465.496 0 2 12094.4851 1 17 2 -13465.496 0 2 12094.4851 1 2 12 2 -22722.174 1 3 -3177.9077 1 3 17 2 -13465.496 0 3 -11772.1338 1 4 17 2 -13465.496 0 3 -415.1114 0 17 2 -3329.619 0 3 -11772.1338 1 6 17 2 -3329.619 0 3 -415.1114 0 7 12 2 -22722.174 1 5 -13311.8247 1 8 15 3 -61284.6900 0 2 16 3 -22905.288 1 3 -61284.6900 0 3 13 3 -12776.884 1 4 28080.1607 0 4 13 3 -12776.884 1 5 -18421.6171 0

If the following commands are issued: rankedF1 <- h2o.rank_within_group_by(train,
c("Group_by_column"), c("Column_to_arrange_by"), c(TRUE), sort_cols_sorted=TRUE)
h2o.summary(rankedF1)
The returned frame will be sorted according to sortCols and hence look like this instead: ID Group_by_column num fdata Column_to_arrange_by num_1
Examples

```r
# Not run:
library(h2o)
h2o.init()

f <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/airlines/allyears2k_headers.zip"
air <- h2o.importFile(f)
group_cols <- c("Distance")
sort_cols <- c("IsArrDelayed", "IsDepDelayed")
sort_directions <- c(TRUE, FALSE)
h2o.rank_within_group_by(x = air, group_by_cols = group_cols,
                          sort_cols = sort_cols,
                          ascending = sort_directions,
                          new_col_name = "New_Rank",
                          sort_cols_sorted = TRUE)
```

```
## End(Not run)
```

### h2o.rapids

**Execute a Rapids expression.**

#### Description

Execute a Rapids expression.

#### Usage

```
h2o.rapids(expr)
```

#### Arguments

- `expr` The rapids expression (ascii string)

#### Examples

```r
# Not run:
h2o.rapids('setproperty "sys.ai.h2o.algos.evaluate_auto_model_parameters" "true"')
```

```
## End(Not run)
```
**h2o.rbind**

*Combine H2O Datasets by Rows*

**Description**

Takes a sequence of H2O data sets and combines them by rows.

**Usage**

```r
h2o.rbind(...)```

**Arguments**

...  
A sequence of H2OFrame arguments. All datasets must exist on the same H2O instance (IP and port) and contain the same number and types of columns.

**Value**

An H2OFrame object containing the combined...arguments row-wise.

**See Also**

`cbind` for the base R method, `rbind()`.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
prostate_rbind <- h2o.rbind(prostate, prostate)
head(prostate_rbind)
dim(prostate)
dim(prostate_rbind)
## End(Not run)
```

---

**h2o.reconstruct**

*Reconstruct Training Data via H2O GLRM Model*

**Description**

Reconstruct the training data and impute missing values from the H2O GLRM model by computing the matrix product of X and Y, and transforming back to the original feature space by minimizing each column’s loss function.
Usage

h2o.reconstruct(object, data, reverse_transform = FALSE)

Arguments

object 
An H2ODimReductionModel object that represents the model to be used for reconstruction.

data 
An H2OFrame object representing the training data for the H2O GLRM model. Used to set the domain of each column in the reconstructed frame.

reverse_transform 
(Optional) A logical value indicating whether to reverse the transformation from model-building by re-scaling columns and adding back the offset to each column of the reconstructed frame.

Value

Returns an H2OFrame object containing the approximate reconstruction of the training data;

See Also

h2o.glrm for making an H2ODimReductionModel.

Examples

## Not run:
library(h2o)
h2o.init()
iris_hf <- as.h2o(iris)
iris_glrm <- h2o.glrm(training_frame = iris_hf, k = 4, transform = "STANDARDIZE",
                    loss = "Quadratic", multi_loss = "Categorical", max_iterations = 1000)
iris_rec <- h2o.reconstruct(iris_glrm, iris_hf, reverse_transform = TRUE)
head(iris_rec)
## End(Not run)

h2o.relevel

Reorders levels of an H2O factor, similarly to standard R's relevel.

Description

The levels of a factor are reordered so that the reference level is at level 0, remaining levels are moved down as needed.

Usage

h2o.relevel(x, y)
Arguments

x  factor column in h2o frame
y  reference level (string)

Value

new reordered factor column

Examples

```r
## Not run:
library(h2o)
h2o.init()

# Convert iris dataset to an H2OFrame
iris_hf <- as.h2o(iris)
# Look at current ordering of the Species column levels
h2o.levels(iris_hf["Species"])
# "setosa" "versicolor" "virginica"
# Change the reference level to "virginica"
iris_hf["Species"] <- h2o.relevel(x = iris_hf["Species"], y = "virginica")
# Observe new ordering
h2o.levels(iris_hf["Species"])
# "virginica" "setosa" "versicolor"

## End(Not run)
```

h2o.removeAll  Remove All Objects on the H2O Cluster

Description

Removes the data from the h2o cluster, but does not remove the local references. Retains models, frames and vectors specified in retained_elements argument. Retained elements must be instances/ids of models and frames only. For models retained, training and validation frames are retained as well. Cross validation models of a retained model are NOT retained automatically, those must be specified explicitly.

Usage

```r
h2o.removeAll(timeout_secs = 0, retained_elements = c())
```

Arguments

timeout_secs  Timeout in seconds. Default is no timeout.
retained_elements  Instances or ids of models and frames to be retained. Combination of instances and ids in the same list is also a valid input.
See Also

h2o.rm

Examples

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
h2o.ls()
h2o.removeAll()
h2o.ls()

## End(Not run)
```

h2o.removeVecs  
*Delete Columns from an H2OFrame*

Description

Delete the specified columns from the H2OFrame. Returns an H2OFrame without the specified columns.

Usage

```
    h2o.removeVecs(data, cols)
```

Arguments

- `data`: The H2OFrame.
- `cols`: The columns to remove.

h2o.rep_len  
*Replicate Elements of Vectors or Lists into H2O*

Description

h2o.rep_len performs just as rep does. It replicates the values in x in the H2O backend.

Usage

```
    h2o.rep_len(x, length.out)
```
h2o.reset_threshold

Reset model threshold and return old threshold value.

**Description**

Reset model threshold and return old threshold value.

**Usage**

```r
h2o.reset_threshold(object, threshold)
```

**Arguments**

- **object**: An `H2OModel` object.
- **threshold**: A threshold value from 0 to 1 included.

**Value**

Returns the previous threshold used in the model.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.importFile(prostate_path)
```

---

**h2o.reset_threshold**

Reset model threshold and return old threshold value.

**Description**

Reset model threshold and return old threshold value.

**Usage**

```r
h2o.reset_threshold(object, threshold)
```

**Arguments**

- **object**: An `H2OFrame`.
- **length.out**: non negative integer. The desired length of the output vector.

**Value**

Creates an `H2OFrame` of the same type as `x`.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

iris <- h2o.importFile(f)
h2o.rep_len(iris, length.out = 3)
```

---

h2o.reset_threshold

Reset model threshold and return old threshold value.

**Description**

Reset model threshold and return old threshold value.

**Usage**

```r
h2o.reset_threshold(object, threshold)
```

**Arguments**

- **object**: An `H2OModel` object.
- **threshold**: A threshold value from 0 to 1 included.

**Value**

Returns the previous threshold used in the model.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.importFile(prostate_path)
```
h2o.residual_analysis_plot

prostate[, 2] <- as.factor(prostate[, 2])
prostate_glm <- h2o.glm(y = "CAPSULE", x = c("AGE", "RACE", "PSA", "DCAPS"),
    training_frame = prostate, family = "binomial",
    nfolds = 0, alpha = 0.5, lambda_search = FALSE)
old_threshold <- h2o.reset_threshold(prostate_glm, 0.9)

## End(Not run)

h2o.residual_analysis_plot

Residual Analysis

Description
Do Residual Analysis and plot the fitted values vs residuals on a test dataset. Ideally, residuals should be randomly distributed. Patterns in this plot can indicate potential problems with the model selection, e.g., using simpler model than necessary, not accounting for heteroscedasticity, autocorrelation, etc. If you notice "striped" lines of residuals, that is just an indication that your response variable was integer valued instead of real valued.

Usage
h2o.residual_analysis_plot(model, newdata)

Arguments

model
An H2OModel.

newdata
An H2OFrame. Used to calculate residuals.

Value
A ggplot2 object

Examples

## Not run:
library(h2o)
h2o.init()

# Import the wine dataset into H2O:
df <- h2o.importFile(f)

# Set the response
test <- "quality"

# Split the dataset into a train and test set:
splits <- h2o.splitFrame(df, ratios = 0.8, seed = 1)
train <- splits[[1]]
test <- splits[[2]]

# Build and train the model:
gbm <- h2o.gbm(y = response,
               training_frame = train)

# Create the residual analysis plot
residual_analysis_plot <- h2o.residual_analysis_plot(gbm, test)
print(residual_analysis_plot)

## End(Not run)

h2o.residual_deviance  Retrieve the residual deviance

**Description**

If "train", "valid", and "xval" parameters are FALSE (default), then the training residual deviance value is returned. If more than one parameter is set to TRUE, then a named vector of residual deviances are returned, where the names are "train", "valid" or "xval".

**Usage**

h2o.residual_deviance(object, train = FALSE, valid = FALSE, xval = FALSE)

**Arguments**

- **object**: An H2OModel or H2OModelMetrics
- **train**: Retrieve the training residual deviance
- **valid**: Retrieve the validation residual deviance
- **xval**: Retrieve the cross-validation residual deviance

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.importFile(prostate_path)
prostate[, 2] <- as.factor(prostate[, 2])
prostate_glm <- h2o.glm(y = "CAPSULE", x = c("AGE", "RACE", "PSA", "DCAPS"),
                        training_frame = prostate, family = "binomial",
                        nfolds = 0, alpha = 0.5, lambda_search = FALSE)
h2o.residual_deviance(prostate_glm, train = TRUE)

## End(Not run)
```
**h2o.residual_dof**

*Retrieve the residual degrees of freedom*

**Description**

If "train", "valid", and "xval" parameters are FALSE (default), then the training residual degrees of freedom value is returned. If more than one parameter is set to TRUE, then a named vector of residual degrees of freedom are returned, where the names are "train", "valid" or "xval".

**Usage**

```r
h2o.residual_dof(object, train = FALSE, valid = FALSE, xval = FALSE)
```

**Arguments**

- `object` - An `H2OModel` or `H2OModelMetrics`
- `train` - Retrieve the training residual degrees of freedom
- `valid` - Retrieve the validation residual degrees of freedom
- `xval` - Retrieve the cross-validation residual degrees of freedom

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.importFile(prostate_path)
prostate[, 2] <- as.factor(prostate[, 2])
prostate_glm <- h2o.glm(y = "CAPSULE", x = c("AGE", "RACE", "PSA", "DCAPS"),
                        training_frame = prostate, family = "binomial",
                        nfolds = 0, alpha = 0.5, lambda_search = FALSE)
h2o.residual_dof(prostate_glm, train = TRUE)
## End(Not run)
```

**h2o.resume**

*Triggers auto-recovery resume - this will look into configured recovery dir and resume and tasks that were interrupted by unexpected cluster stopping.*

**Description**

Triggers auto-recovery resume - this will look into configured recovery dir and resume and tasks that were interrupted by unexpected cluster stopping.
**Usage**
```
h2o.resume(recovery_dir = NULL)
```

**Arguments**
- **recovery_dir**: A character path to where cluster recovery data is stored, if blank, will use cluster's configuration.

**h2o.resumeGrid**
Resume previously stopped grid training.

**Description**
Resume previously stopped grid training.

**Usage**
```
h2o.resumeGrid(grid_id, recovery_dir = NULL, ...)
```

**Arguments**
- **grid_id**: ID of existing grid search
- **recovery_dir**: When specified the grid and all necessary data (frames, models) will be saved to this directory (use HDFS or other distributed file-system). Should the cluster crash during training, the grid can be reloaded from this directory via h2o.loadGrid and training can be resumed
- **...**: Additional parameters to modify the resumed Grid.

**h2o.rm**
Delete Objects In H2O

**Description**
Remove the h2o Big Data object(s) having the key name(s) from ids.

**Usage**
```
h2o.rm(ids, cascade = TRUE)
```

**Arguments**
- **ids**: The object or hex key associated with the object to be removed or a vector/list of those things.
- **cascade**: Boolean, if set to TRUE (default), the object dependencies (e.g. submodels) are also removed.
**h2o.rmse**

**See Also**

h2o.assign, h2o.ls

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
iris <- as.h2o(iris)
model <- h2o.glm(1:4,5,training = iris, family = "multinomial")
h2o.rm(iris)
## End(Not run)
```

---

**h2o.rmse**

Retrieves Root Mean Squared Error Value

**Description**

Retrieves the root mean squared error value from an H2OModelMetrics object. If "train", "valid", and "xval" parameters are FALSE (default), then the training RMSE value is returned. If more than one parameter is set to TRUE, then a named vector of RMSEs are returned, where the names are "train", "valid" or "xval".

**Usage**

```r
h2o.rmse(object, train = FALSE, valid = FALSE, xval = FALSE)
```

**Arguments**

- `object`: An H2OModelMetrics object of the correct type.
- `train`: Retrieve the training RMSE
- `valid`: Retrieve the validation RMSE
- `xval`: Retrieve the cross-validation RMSE

**Details**

This function only supports H2OBinomialMetrics, H2OMultinomialMetrics, and H2ORegressionMetrics objects.

**See Also**

h2o.auc for AUC, h2o.mse for RMSE, and h2o.metric for the various threshold metrics. See h2o.performance for creating H2OModelMetrics objects.
Examples

```r
## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(prostate_path)

prostate[, 2] <- as.factor(prostate[, 2])
model <- h2o.gbm(x = 3:9, y = 2, training_frame = prostate, distribution = "bernoulli")
perf <- h2o.performance(model, prostate)
h2o.rmse(perf)

## End(Not run)
```

---

**h2o.rmsle**

*Retrieve the Root Mean Squared Log Error*

### Description

Retrieves the root mean squared log error (RMSLE) value from an H2O model. If "train", "valid", and "xval" parameters are FALSE (default), then the training rmsle value is returned. If more than one parameter is set to TRUE, then a named vector of rmsles are returned, where the names are "train", "valid" or "xval".

### Usage

```r
h2o.rmsle(object, train = FALSE, valid = FALSE, xval = FALSE)
```

### Arguments

- `object`: An `H2OModel` object.
- `train`: Retrieve the training rmsle
- `valid`: Retrieve the validation set rmsle if a validation set was passed in during model build time.
- `xval`: Retrieve the cross-validation rmsle

### Examples

```r
## Not run:
library(h2o)

h <- h2o.init()
fr <- as.h2o(iris)
m <- h2o.deeplearning(x = 2:5, y = 1, training_frame = fr)
h2o.rmsle(m)
```
h2o.round

## End(Not run)

---

**h2o.round**  
*Round doubles/floats to the given number of decimal places.*

### Description

Round doubles/floats to the given number of decimal places.

### Usage

```r
h2o.round(x, digits = 0)
```

```r
round(x, digits = 0)
```

### Arguments

- **x**: An H2OFrame object.
- **digits**: Number of decimal places to round doubles/floats. Rounding to a negative number of decimal places is

### See Also

`Round` for the base R implementation, `round()`.

### Examples

```r
## Not run:
library(h2o)
h2o.init()

f <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/coxph_test/heart.csv"
heart <- h2o.importFile(f)

h2o.round(heart["age"], digits = 3)
```

## End(Not run)
h2o.rstrip  

Strip set from right

Description

Return a copy of the target column with trailing characters removed. The set argument is a string specifying the set of characters to be removed. If omitted, the set argument defaults to removing whitespace.

Usage

h2o.rstrip(x, set = " ")

Arguments

x  
The column whose strings should be rstrip-ed.

set  
string of characters to be removed

Examples

## Not run:
library(h2o)
h2o.init()
string_to_rstrip <- as.h2o("1234567890")
rstrip_string <- h2o.rstrip(string_to_rstrip, "890") #Remove "890"

## End(Not run)

h2o.rulefit  
Build a RuleFit Model

Description

Builds a Distributed RuleFit model on a parsed dataset, for regression or classification.

Usage

h2o.rulefit(
  x,
  y,
  training_frame,
  model_id = NULL,
  validation_frame = NULL,
  seed = -1,
  algorithm = c("AUTO", "DRF", "GBM"),
  min_rule_length = 3,
Arguments

x  (Optional) A vector containing the names or indices of the predictor variables to use in building the model. If x is missing, then all columns except y are used.
y  The name or column index of the response variable in the data. The response must be either a numeric or a categorical/factor variable. If the response is numeric, then a regression model will be trained, otherwise it will train a classification model.

training_frame  Id of the training data frame.
model_id  Destination id for this model; auto-generated if not specified.
validation_frame  Id of the validation data frame.
seed  Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).

algorithm  The algorithm to use to generate rules. Must be one of: "AUTO", "DRF", "GBM". Defaults to AUTO.

min_rule_length  Minimum length of rules. Defaults to 3.
max_rule_length  Maximum length of rules. Defaults to 3.

max_num_rules  The maximum number of rules to return. defaults to -1 which means the number of rules is selected by diminishing returns in model deviance. Defaults to -1.
model_type  Specifies type of base learners in the ensemble. Must be one of: "rules_and_linear", "rules", "linear". Defaults to rules_and_linear.
weights_column  Column with observation weights. Giving some observation a weight of zero is equivalent to excluding it from the dataset; giving an observation a relative weight of 2 is equivalent to repeating that row twice. Negative weights are not allowed. Note: Weights are per-row observation weights and do not increase the size of the data frame. This is typically the number of times a row is repeated, but non-integer values are supported as well. During training, rows with higher weights matter more, due to the larger loss function pre-factor. If you set weight = 0 for a row, the returned prediction frame at that row is zero and this is incorrect. To get an accurate prediction, remove all rows with weight == 0.
**distribution**

Distribution function. Must be one of: "AUTO", "bernoulli", "multinomial", "gaussian", "poisson", "gamma", "tweedie", "laplace", "quantile", "huber". Defaults to AUTO.

**rule_generation_ntrees**

Specifies the number of trees to build in the tree model. Defaults to 50. Defaults to 50.

**auc_type**

Set default multinomial AUC type. Must be one of: "AUTO", "NONE", "MACRO_OVR", "WEIGHTED_OVR", "MACRO_OVO", "WEIGHTED_OVO". Defaults to AUTO.

**remove_duplicates**

Logical. Whether to remove rules which are identical to an earlier rule. Defaults to true. Defaults to TRUE.

**lambda**

Lambda for LASSO regressor.

---

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

# Import the titanic dataset:
f <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/gbm_test/titanic.csv"
coltypes <- list(by.col.name = c("pclass", "survived"), types=c("Enum", "Enum"))
df <- h2o.importFile(f, col.types = coltypes)

# Split the dataset into train and test
splits <- h2o.splitFrame(data = df, ratios = 0.8, seed = 1)
train <- splits[[1]]
test <- splits[[2]]

# Set the predictors and response; set the factors:
response <- "survived"
predictors <- c("age", "sibsp", "parch", "fare", "sex", "pclass")

# Build and train the model:
rfit <- h2o.rulefit(y = response,
                   x = predictors,
                   training_frame = train,
                   max_rule_length = 10,
                   max_num_rules = 100,
                   seed = 1)

# Retrieve the rule importance:
print(rfit$model$rule_importance)

# Predict on the test data:
h2o.predict(rfit, newdata = test)

## End(Not run)
```
### h2o.runif

**Produce a Vector of Random Uniform Numbers**

**Description**

Creates a vector of random uniform numbers equal in length to the length of the specified H2O dataset.

**Usage**

```r
h2o.runif(x, seed = -1)
```

**Arguments**

- `x`: An H2OFrame object.
- `seed`: A random seed used to generate draws from the uniform distribution.

**Value**

A vector of random, uniformly distributed numbers. The elements are between 0 and 1.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.importFile(path = prostate_path)
s <- h2o.runif(prostate)
summary(s)
prostate_train <- prostate[s <= 0.8,]
prostate_test <- prostate[s > 0.8,]
nrow(prostate_train) + nrow(prostate_test)

## End(Not run)
```

### h2o.saveGrid

**Saves an existing Grid of models into a given folder.**

**Description**

Returns a reference to the saved Grid.
h2o.saveGrid

Usage

h2o.saveGrid(
  grid_directory,
  grid_id,
  save_params_references = FALSE,
  export_cross_validation_predictions = FALSE
)

Arguments

grid_directory  A character string containing the path to the folder for the grid to be saved to.
grid_id         A character string with identification of the grid to be saved.
save_params_references
    A logical indicating if objects referenced by grid parameters (e.g. training frame, calibration frame) should also be saved.
export_cross_validation_predictions
    A logical indicating whether exported model artifacts should also include CV holdout Frame predictions.

Value

Returns an object that is a subclass of H2OGrid.

Examples

## Not run:
library(h2o)
h2o.init()

iris <- as.h2o(iris)

ntrees_opts = c(1, 5)
learn_rate_opts = c(0.1, 0.01)
size_of_hyper_space = length(ntrees_opts) * length(learn_rate_opts)

hyper_parameters = list(ntrees = ntrees_opts, learn_rate = learn_rate_opts)
# Tempdir is chosen arbitrarily. May be any valid folder on an H2O-supported filesystem.
baseline_grid <- h2o.grid(algorithm = "gbm",
                          grid_id = "gbm_grid_test",
                          x = 1:4,
                          y = 5,
                          training_frame = iris,
                          hyper_params = hyper_parameters)

grid_path <- h2o.saveGrid(grid_directory = tempdir(), grid_id = baseline_grid@grid_id)
# Remove everything from the cluster or restart it
h2o.removeAll()
grid <- h2o.loadGrid(grid_path)

## End(Not run)
h2o.saveModel

Save an H2O Model Object to Disk

Description

Save an H2OModel to disk. (Note that ensemble binary models can be saved.)

Usage

h2o.saveModel(
  object,
  path = "",
  force = FALSE,
  export_cross_validation_predictions = FALSE,
  filename = ""
)

Arguments

object      an H2OModel object.
path        string indicating the directory the model will be written to.
force       logical, indicates how to deal with files that already exist.
export_cross_validation_predictions logical, indicates whether the exported model artifacts should also include CV Holdout Frame predictions. Default is not to export the predictions.
filename    string indicating the file name.

Details

In the case of existing files force = TRUE will overwrite the file. Otherwise, the operation will fail. The owner of the file saved is the user by which H2O cluster was executed.

See Also

h2o.loadModel for loading a model to H2O from disk

Examples

## Not run:
# library(h2o)
# h2o.init()
# prostate <- h2o.importFile(path = paste("https://raw.githubusercontent.com/",
#     "h2oai/h2o-3/master/smalldata/logreg/prostate.csv", sep = "/"))
# prostate_glm <- h2o.glm(y = "CAPSULE", x = c("AGE", "RACE", "PSA", "DCAPS"),
#     training_frame = prostate, family = "binomial", alpha = 0.5)
# h2o.saveModel(object = prostate_glm, path = "/Users/UserName/Desktop", force = TRUE)

## End(Not run)
h2o.saveModelDetails  
Save an H2O Model Details

Description
Save Model Details of an H2O Model in JSON Format

Usage
h2o.saveModelDetails(object, path = "", force = FALSE, filename = "")

Arguments
- object: an H2OModel object.
- path: string indicating the directory the model details will be written to.
- force: logical, indicates how to deal with files that already exist.
- filename: string indicating the file name. (Type of file is always .json)

Details
Model Details will download as a JSON file. In the case of existing files force = TRUE will overwrite the file. Otherwise, the operation will fail.

Examples
```r
## Not run:
# library(h2o)
# h2o.init()
# prostate <- h2o.uploadFile(path = system.file("extdata", "prostate.csv", package = "h2o"))
# prostate_glm <- h2o.glm(y = "CAPSULE", x = c("AGE", "RACE", "PSA", "DCAPS"),
#                         training_frame = prostate, family = "binomial", alpha = 0.5)
# h2o.saveModelDetails(object = prostate_glm, path = "/Users/UserName/Desktop", force = TRUE)
## End(Not run)
```

h2o.saveMojo  
Deprecated - use h2o.save_mojo instead. Save an H2O Model Object as Mojo to Disk

Description
Save an MOJO (Model Object, Optimized) to disk.

Usage
h2o.saveMojo(object, path = "", force = FALSE)
**h2o.save_frame**

**Description**

Store frame data in H2O’s native format.

**Usage**

```r
h2o.save_frame(x, dir, force = TRUE)
```

**Arguments**

- **x**: An H2OFrame object
- **dir**: a filesystem location where to write frame data (hdfs, nfs)
- **force**: logical. overwrite already existing files (defaults to true)

**Arguments**

- **object**: an H2OModel object.
- **path**: string indicating the directory the model will be written to.
- **force**: logical, indicates how to deal with files that already exist.

**Details**

MOJO will download as a zip file. In the case of existing files force = TRUE will overwrite the file. Otherwise, the operation will fail.

**See Also**

`h2o.saveModel` for saving a model to disk as a binary object.

**Examples**

```r
## Not run:
# library(h2o)
# h2o.init()
# prostate <- h2o.uploadFile(path = system.file("extdata", "prostate.csv", package="h2o"))
# prostate_glm <- h2o.glm(y = "CAPSULE", x = c("AGE", "RACE", "PSA", "DCAPS"),
#                        training_frame = prostate, family = "binomial", alpha = 0.5)
# h2o.saveMojo(object = prostate_glm, path = "/Users/UserName/Desktop", force = TRUE)
## End(Not run)
```
Examples

```r
## Not run:
library(h2o)
h2o.init()

prostate_path = system.file("extdata", "prostate.csv", package = "h2o")
prostate = h2o.importFile(path = prostate_path)
h2o.save_frame(prostate, "/tmp/prostate")

## End(Not run)
```

---

**h2o.save_mojo**

Save an H2O Model Object as Mojo to Disk

Description

Save an MOJO (Model Object, Optimized) to disk.

Usage

```r
h2o.save_mojo(object, path = "", force = FALSE, filename = "")
```

Arguments

- `object`: an `H2OModel` object.
- `path`: string indicating the directory the model will be written to.
- `force`: logical, indicates how to deal with files that already exist.
- `filename`: string indicating the file name. (Type of file is always .zip)

Details

MOJO will download as a zip file. In the case of existing files `force = TRUE` will overwrite the file. Otherwise, the operation will fail.

See Also

- `h2o.saveModel` for saving a model to disk as a binary object.

Examples

```r
## Not run:
# library(h2o)
# h2o.init()
# prostate <- h2o.uploadFile(path = system.file("extdata", "prostate.csv", package="h2o"))
# prostate_glm <- h2o.glm(y = "CAPSULE", x = c("AGE", "RACE", "PSA", "DCAPS"),
# training_frame = prostate, family = "binomial", alpha = 0.5)
# h2o.save_mojo(object = prostate_glm, path = "/Users/UserName/Desktop", force = TRUE)

## End(Not run)
```
**h2o.save_to_hive**  
*Save contents of this data frame into a Hive table*

**Description**

For example, h2o.save_to_hive(data_frame, "jdbc:hive2://host:10000/database", "table_name") h2o.save_to_hive(data_frame, "jdbc:hive2://host:10000/", "database.table_name", format = "parquet")

**Usage**

h2o.save_to_hive(
  data,
  jdbc_url,
  table_name,
  format = "csv",
  table_path = NULL,
  tmp_path = NULL
)

**Arguments**

data: A H2O Frame object to be saved.
jdbc_url: Hive JDBC connection URL.
table_name: Table name into which to store the data. The table must not exist as it will be created.
format: Storage format of created Hive table. (default csv, can be csv or parquet)
table_path: If specified, the table will be created as an external table and this is where the data

tmp_path: Path where to store temporary data.

**h2o.scale**  
*Scaling and Centering of an H2OFrame*

**Description**

Centers and/or scales the columns of an H2O dataset.

**Usage**

h2o.scale(x, center = TRUE, scale = TRUE, inplace = FALSE)
Arguments

x  An H2OFrame object.
center  either a logical value or numeric vector of length equal to the number of columns of x.
scale  either a logical value or numeric vector of length equal to the number of columns of x.
inplace  a logical values indicating whether directly overwrite original data (disabled by default). Exposed for backwards compatibility (prior versions of this functions were always doing an inplace update).

Examples

## Not run:
library(h2o)
h2o.init()
iris_hf <- as.h2o(iris)
summary(iris_hf)

# Scale and center all the numeric columns in iris data set
iris_scaled <- h2o.scale(iris_hf[, 1:4])

## End(Not run)

h2o.scoreHistory  Retrieve Model Score History

Description

Retrieve Model Score History

Usage

h2o.scoreHistory(object)

Arguments

object  An H2OModel object.

Examples

## Not run:
library(h2o)
h2o.init()

cars <- h2o.importFile(f)
cars[economy_20mpg] <- as.factor(cars[economy_20mpg])
predictors <- c("displacement", "power", "weight", "acceleration", "year")
response <- "economy_20mpg"
cars_split <- h2o.splitFrame(data = cars, ratios = 0.8, seed = 1234)
train <- cars_split[[1]]
valid <- cars_split[[2]]
cars_gbm <- h2o.gbm(x = predictors, y = response,
                   training_frame = train,
                   validation_frame = valid,
                   seed = 1234)
h2o.scoreHistory(cars_gbm)

## End(Not run)

h2o.scoreHistoryGAM  

Retrieve GLM Model Score History buried in GAM model

Description
Retrieve GLM Model Score History buried in GAM model

Usage
h2o.scoreHistoryGAM(object)

Arguments
object  

An H2OModel object.

h2o.screeplot  

Scree Plot

Description
Scree Plot

Usage
h2o.screeplot(model, type = c("barplot", "lines"))

Arguments
model  

A PCA model

type  

Type of the plot. Either "barplot" or "lines".
h2o.sdev

**Description**

Retrieve the standard deviations of principal components

**Usage**

h2o.sdev(object)

---

h2o.sd

*Standard Deviation of a column of data.*

**Description**

Obtain the standard deviation of a column of data.

**Usage**

h2o.sd(x, na.rm = FALSE)

sd(x, na.rm = FALSE)

**Arguments**

x

An H2OFrame object.

na.rm

logical. Should missing values be removed?

**See Also**

h2o.var for variance, and sd for the base R implementation.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
sd(prostate$AGE)
## End(Not run)
```
Arguments

object      An H2ODimReductionModel object.

Examples

```r
## Not run:
library(h2o)
h2o.init()

cars <- h2o.importFile(f)
predictors <- c("displacement", "power", "weight", "acceleration", "year")
cars_pca <- h2o.prcomp(cars, transform = "STANDARDIZE",
                        k = 3, x = predictors, seed = 12345)
h2o.sdev(cars_pca)

## End(Not run)
```

---

h2o.setLevels       Set Levels of H2O Factor Column

Description

Works on a single categorical vector. New domains must be aligned with the old domains. This call has SIDE EFFECTS and mutates the column in place (change of the levels will also affect all the frames that are referencing this column). If you want to make a copy of the column instead, use parameter in.place = FALSE.

Usage

h2o.setLevels(x, levels, in.place = TRUE)

Arguments

- x      A single categorical column.
- levels A character vector specifying the new levels. The number of new levels must match the number of old levels.
- in.place Indicates whether new domain will be directly applied to the column (in place change) or if a copy of the column will be created with the given domain levels.

Examples

```r
## Not run:
library(h2o)
h2o.init()

iris_hf <- as.h2o(iris)
new_levels <- c("setosa", "versicolor", "caroliniana")
```
iris_hf$Species <- h2o.setLevels(iris_hf$Species, new_levels, in.place = FALSE)
h2o.levels(iris_hf$Species)

## End(Not run)

### h2o.setTimezone

Set the Time Zone on the H2O cluster

**Description**

Set the Time Zone on the H2O cluster

**Usage**

h2o.setTimezone(tz)

**Arguments**

- **tz**
  
The desired timezone.

**Examples**

## Not run:
library(h2o)
h2o.init()
h2o.setTimezone("America/Juneau")
h2o.getTimezone()

## End(Not run)

### h2o.set_s3_credentials

Creates a new Amazon S3 client internally with specified credentials.

**Description**

There are no validations done to the credentials. Incorrect credentials are thus revealed with first S3 import call.

**Usage**

h2o.set_s3_credentials(secretKeyId, secretAccessKey, sessionToken = NULL)
Arguments

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>secretKeyId</td>
<td>Amazon S3 Secret Key ID (provided by Amazon)</td>
</tr>
<tr>
<td>secretAccessKey</td>
<td>Amazon S3 Secret Access Key (provided by Amazon)</td>
</tr>
<tr>
<td>sessionToken</td>
<td>Amazon Session Token (optional, only when using AWS Temporary Credentials)</td>
</tr>
</tbody>
</table>

Description

SHAP explanation shows contribution of features for a given instance. The sum of the feature contributions and the bias term is equal to the raw prediction of the model, i.e., prediction before applying inverse link function. H2O implements TreeSHAP which when the features are correlated, can increase contribution of a feature that had no influence on the prediction.

Usage

```r
h2o.shap_explain_row_plot(
  model, 
  newdata, 
  row_index, 
  columns = NULL, 
  top_n_features = 10, 
  plot_type = c("barplot", "breakdown"), 
  contribution_type = c("both", "positive", "negative")
)
```

Arguments

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>model</td>
<td>An H2O tree-based model. This includes Random Forest, GBM and XGboost only. Must be a binary classification or regression model.</td>
</tr>
<tr>
<td>newdata</td>
<td>An H2O Frame, used to determine feature contributions.</td>
</tr>
<tr>
<td>row_index</td>
<td>Instance row index.</td>
</tr>
<tr>
<td>columns</td>
<td>List of columns or list of indices of columns to show. If specified, then the top_n_features parameter will be ignored.</td>
</tr>
<tr>
<td>top_n_features</td>
<td>Integer specifying the maximum number of columns to show (ranked by their contributions). When plot_type = &quot;barplot&quot;, then top_n_features features will be chosen for each contribution_type.</td>
</tr>
<tr>
<td>plot_type</td>
<td>Either &quot;barplot&quot; or &quot;breakdown&quot;. Defaults to &quot;barplot&quot;.</td>
</tr>
<tr>
<td>contribution_type</td>
<td>When plot_type == &quot;barplot&quot;, plot one of &quot;negative&quot;, &quot;positive&quot;, or &quot;both&quot; contributions. Defaults to &quot;both&quot;.</td>
</tr>
</tbody>
</table>
h2o.shap_summary_plot

Value

A ggplot2 object.

Examples

```r
## Not run:
library(h2o)
h2o.init()

# Import the wine dataset into H2O:
df <- h2o.importFile(f)

# Set the response
response <- "quality"

# Split the dataset into a train and test set:
splits <- h2o.splitFrame(df, ratios = 0.8, seed = 1)
train <- splits[[1]]
test <- splits[[2]]

# Build and train the model:
gbm <- h2o.gbm(y = response,
        training_frame = train)

# Create the SHAP row explanation plot
shap_explain_row_plot <- h2o.shap_explain_row_plot(gbm, test, row_index = 1)
print(shap_explain_row_plot)

## End(Not run)
```

h2o.shap_summary_plot  SHAP Summary Plot

Description

SHAP summary plot shows the contribution of the features for each instance (row of data). The sum of the feature contributions and the bias term is equal to the raw prediction of the model, i.e., prediction before applying inverse link function.

Usage

```r
h2o.shap_summary_plot(
    model,
    newdata,
    columns = NULL,
    top_n_features = 20,
    sample_size = 1000
)
```
Arguments

- **model**: An H2O tree-based model. This includes Random Forest, GBM and XGboost only. Must be a binary classification or regression model.
- **newdata**: An H2O Frame, used to determine feature contributions.
- **columns**: List of columns or list of indices of columns to show. If specified, then the `top_n_features` parameter will be ignored.
- **top_n_features**: Integer specifying the maximum number of columns to show (ranked by variable importance).
- **sample_size**: Integer specifying the maximum number of observations to be plotted.

Value

A ggplot2 object

Examples

```r
## Not run:
library(h2o)
h2o.init()

# Import the wine dataset into H2O:
df <- h2o.importFile(f)

# Set the response
test <- "quality"

# Split the dataset into a train and test set:
splits <- h2o.splitFrame(df, ratios = 0.8, seed = 1)
train <- splits[[1]]
test <- splits[[2]]

# Build and train the model:
gbm <- h2o.gbm(y = response,
               training_frame = train)

# Create the SHAP summary plot
shap_summary_plot <- h2o.shap_summary_plot(gbm, test)
print(shap_summary_plot)

## End(Not run)
```

Description

Enable Progress Bar
h2o.shutdown

Usage

h2o.show_progress()

Examples

## Not run:
library(h2o)
h2o.init()
h2o.no_progress()

iris <- h2o.importFile(f)
iris["class"] <- as.factor(iris["class"])
predictors <- c("sepal_len", "sepal_wid", "petal_len", "petal_wid")
splits <- h2o.splitFrame(iris, ratios = 0.8, seed = 1234)
train <- splits[[1]]
valid <- splits[[2]]
h2o.show_progress()

iris_km <- h2o.kmeans(x = predictors,
                       training_frame = train,
                       validation_frame = valid,
                       k = 10, estimate_k = TRUE,
                       standardize = FALSE, seed = 1234)

## End(Not run)

---

h2o.shutdown

Shut Down H2O Instance

Description

Shut down the specified instance. All data will be lost.

Usage

h2o.shutdown(prompt = TRUE)

Arguments

prompt A logical value indicating whether to prompt the user before shutting down the H2O server.

Details

This method checks if H2O is running at the specified IP address and port, and if it is, shuts down that H2O instance.
WARNING

All data, models, and other values stored on the server will be lost! Only call this function if you and all other clients connected to the H2O server are finished and have saved your work.

Note

Users must call h2o.shutdown explicitly in order to shut down the local H2O instance started by R. If R is closed before H2O, then an attempt will be made to automatically shut down H2O. This only applies to local instances started with h2o.init, not remote H2O servers.

See Also

h2o.init

Examples

# Don't run automatically to prevent accidentally shutting down a cluster
## Not run:
library(h2o)
h2o.init()
h2o.shutdown()

## End(Not run)

h2o.signif

Round doubles/floats to the given number of significant digits.

Description

Round doubles/floats to the given number of significant digits.

Usage

h2o.signif(x, digits = 6)

signif(x, digits = 6)

Arguments

x          An H2OFrame object.
digits     Number of significant digits to round doubles/floats.

See Also

Round for the base R implementation, signif().
h2o.sin

Examples

## Not run:
library(h2o)
h2o.init()

f <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/coxph_test/heart.csv"
heart <- h2o.importFile(f)

h2o.signif(heart["age"], digits = 3)

## End(Not run)

h2o.sin

Compute the sine of x

Description

Compute the sine of x

Usage

h2o.sin(x)

Arguments

x An H2OFrame object.

See Also

`Trig` for the base R implementation, `sin()`.

Examples

## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)

h2o.sin(frame)

## End(Not run)
h2o.skewness

Skewness of a column

Description
Obtain the skewness of a column of a parsed H2O data object.

Usage
h2o.skewness(x, ..., na.rm = TRUE)
skewness.H2OFrame(x, ..., na.rm = TRUE)

Arguments
x An H2OFrame object.
... Further arguments to be passed from or to other methods.
na.rm A logical value indicating whether NA or missing values should be stripped before the computation.

Value
Returns a list containing the skewness for each column (NaN for non-numeric columns).

Examples
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
h2o.skewness(prostate$AGE)
## End(Not run)

h2o.splitFrame
Split an H2O Data Set

Description
Split an existing H2O data set according to user-specified ratios. The number of subsets is always 1 more than the number of given ratios. Note that this does not give an exact split. H2O is designed to be efficient on big data using a probabilistic splitting method rather than an exact split. For example, when specifying a split of 0.75/0.25, H2O will produce a test/train split with an expected value of 0.75/0.25 rather than exactly 0.75/0.25. On small datasets, the sizes of the resulting splits will deviate from the expected value more than on big data, where they will be very close to exact.
Usage

h2o.splitFrame(data, ratios = 0.75, destination_frames, seed = -1)

Arguments

data An H2OFrame object, to be split.
ratios A numeric value or array indicating the ratio of total rows contained in each split. Must total up to less than 1. e.g. c(0.8) for 80/20 split.
destination_frames An array of frame IDs equal to the number of values specified in the ratios array, plus one.
seed Random seed.

Value

Returns a list of split H2OFrames

Examples

## Not run:
library(h2o)
h2o.init()
iris_hf <- as.h2o(iris)
iris_split <- h2o.splitFrame(iris_hf, ratios = c(0.2, 0.5))
head(iris_split[[1]])
summary(iris_split[[1]])

## End(Not run)

h2o.sqrt

Compute the square root of x

Description

Compute the square root of x

Usage

h2o.sqrt(x)

Arguments

x An H2OFrame object.

See Also

MathFun for the base R implementation, sqrt().
Examples

```r
## Not run:
library(h2o)
#h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)

h2o.sqrt(frame)

## End(Not run)
```

h2o.stackedEnsemble  
Builds a Stacked Ensemble

Description

Build a stacked ensemble (aka. Super Learner) using the H2O base learning algorithms specified by the user.

Usage

```r
h2o.stackedEnsemble(
  x,
  y,
  training_frame,
  model_id = NULL,
  validation_frame = NULL,
  blending_frame = NULL,
  base_models = list(),
  metalearner_algorithm = c("AUTO", "deeplearning", "drf", "gbm", "glm", "naivebayes", "xgboost"),
  metalearner_nfolds = 0,
  metalearner_fold_assignment = c("AUTO", "Random", "Modulo", "Stratified"),
  metalearner_fold_column = NULL,
  metalearner_params = NULL,
  metalearner_transform = c("NONE", "Logit"),
  max_runtime_secs = 0,
  weights_column = NULL,
  offset_column = NULL,
  seed = -1,
  score_training_samples = 10000,
  keep_levelone_frame = FALSE,
  export_checkpoints_dir = NULL,
  auc_type = c("AUTO", "NONE", "MACRO_OVR", "WEIGHTED_OVR", "MACRO_OVO", "WEIGHTED_OVO")
)
```
**Arguments**

- **x** *(Optional).* A vector containing the names or indices of the predictor variables to use in building the model. If x is missing, then all columns except y are used. Training frame is used only to compute ensemble training metrics.

- **y** The name or column index of the response variable in the data. The response must be either a numeric or a categorical/factor variable. If the response is numeric, then a regression model will be trained, otherwise it will train a classification model.

- **training_frame** Id of the training data frame.

- **model_id** Destination id for this model; auto-generated if not specified.

- **validation_frame** Id of the validation data frame.

- **blending_frame** Frame used to compute the predictions that serve as the training frame for the metalearner (triggers blending mode if provided)

- **base_models** List of models or grids (or their ids) to ensemble/stack together. Grids are expanded to individual models. If not using blending frame, then models must have been cross-validated using nfolds > 1, and folds must be identical across models.

- **metalearner_algorithm** Type of algorithm to use as the metalearner. Options include 'AUTO' (GLM with non negative weights; if validation_frame is present, a lambda search is performed), 'deeplearning' (Deep Learning with default parameters), 'drf' (Random Forest with default parameters), 'gbm' (GBM with default parameters), 'glm' (GLM with default parameters), 'naivebayes' (NaiveBayes with default parameters), or 'xgboost' (if available, XGBoost with default parameters). Must be one of: "AUTO", "deeplearning", "drf", "gbm", "glm", "naivebayes", "xgboost". Defaults to AUTO.

- **metalearner_nfolds** Number of folds for K-fold cross-validation of the metalearner algorithm (0 to disable or >= 2). Defaults to 0.

- **metalearner_fold_assignment** Cross-validation fold assignment scheme for metalearner cross-validation. Defaults to AUTO (which is currently set to Random). The 'Stratified' option will stratify the folds based on the response variable, for classification problems. Must be one of: "AUTO", "Random", "Modulo", "Stratified".

- **metalearner_fold_column** Column with cross-validation fold index assignment per observation for cross-validation of the metalearner.

- **metalearner_params** Parameters for metalearner algorithm

- **metalearner_transform** Transformation used for the level one frame. Must be one of: "NONE", "Logit". Defaults to NONE.

- **max_runtime_secs** Maximum allowed runtime in seconds for model training. Use 0 to disable. Defaults to 0.
weights_column  Column with observation weights. Giving some observation a weight of zero is equivalent to excluding it from the dataset; giving an observation a relative weight of 2 is equivalent to repeating that row twice. Negative weights are not allowed. Note: Weights are per-row observation weights and do not increase the size of the data frame. This is typically the number of times a row is repeated, but non-integer values are supported as well. During training, rows with higher weights matter more, due to the larger loss function pre-factor. If you set weight = 0 for a row, the returned prediction frame at that row is zero and this is incorrect. To get an accurate prediction, remove all rows with weight == 0.

offset_column  Offset column. This will be added to the combination of columns before applying the link function.

seed  Seed for random numbers; passed through to the metalearner algorithm. Defaults to -1 (time-based random number).

score_training_samples  Specify the number of training set samples for scoring. The value must be >= 0. To use all training samples, enter 0. Defaults to 10000.

keep_levelone_frame  Logical. Keep level one frame used for metalearner training. Defaults to FALSE.

export_checkpoints_dir  Automatically export generated models to this directory.

auc_type  Set default multinomial AUC type. Must be one of: "AUTO", "NONE", "MACRO_OVR", "WEIGHTED_OVR", "MACRO_OVO", "WEIGHTED_OVO". Defaults to AUTO.

Examples

## Not run:
library(h2o)
h2o.init()

# Import a sample binary outcome train/test set
train <- h2o.importFile("https://s3.amazonaws.com/erin-data/higgs/higgs_train_10k.csv")
test <- h2o.importFile("https://s3.amazonaws.com/erin-data/higgs/higgs_test_5k.csv")

# Identify predictors and response
y <- "response"
x <- setdiff(names(train), y)

# For binary classification, response should be a factor
train[, y] <- as.factor(train[, y])
test[, y] <- as.factor(test[, y])

# Number of CV folds
nfolds <- 5

# Train & Cross-validate a GBM
my_gbm <- h2o.gbm(x = x,
                  y = y,
                  training_frame = train,
distribution = "bernoulli",
ntrees = 10,
max_depth = 3,
min_rows = 2,
learn_rate = 0.2,
nfolds = nfolds,
fold_assignment = "Modulo",
keep_cross_validation_predictions = TRUE,
seed = 1)

# Train & Cross-validate a RF
my_rf <- h2o.randomForest(x = x,
    y = y,
    training_frame = train,
    ntree = 50,
    nfolds = nfolds,
    fold_assignment = "Modulo",
    keep_cross_validation_predictions = TRUE,
    seed = 1)

# Train a stacked ensemble using the GBM and RF above
ensemble <- h2o.stackedEnsemble(x = x,
    y = y,
    training_frame = train,
    model_id = "my_ensemble_binomial",
    base_models = list(my_gbm, my_rf))

## End(Not run)

---

### h2o.startLogging

**Start Writing H2O R Logs**

**Description**

Begin logging H2O R POST commands and error responses to local disk. Used primarily for debugging purposes.

**Usage**

h2o.startLogging(file)

**Arguments**

- **file** a character string name for the file, automatically generated

**See Also**

h2o.stopLogging,h2o.clearLog,h2o.openLog
## Not run:
library(h2o)
h2o.init()
h2o.startLogging()
australia_path = system.file("extdata", "australia.csv", package = "h2o")
australia = h2o.importFile(path = australia_path)
h2o.stopLogging()

## End(Not run)

### Description
Plot a GLM model’s standardized coefficient magnitudes.

### Usage

```r
h2o.std_coef_plot(model, num_of_features = NULL)
```

### Arguments

- **model**: A trained generalized linear model
- **num_of_features**: The number of features to be shown in the plot

### See Also

- `h2o.varimp_plot` for variable importances plot of random forest, GBM, deep learning.

### Examples

## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.importFile(prostate_path)
prostate[, 2] <- as.factor(prostate[, 2])
prostate_glm <- h2o.glm(y = "CAPSULE", x = c("AGE", "RACE", "PSA", "DCAPS"),
training_frame = prostate, family = "binomial",
nfolds = 0, alpha = 0.5, lambda_search = FALSE)
h2o.std_coef_plot(prostate_glm)

## End(Not run)
**h2o.stopLogging**  
*Stop Writing H2O R Logs*

**Description**

Halt logging of H2O R POST commands and error responses to local disk. Used primarily for debugging purposes.

**Usage**

```r
eh2o.stopLogging()
```

**See Also**

`h2o.startLogging, h2o.clearLog, h2o.openLog`

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
h2o.startLogging()
australia_path = system.file("extdata", "australia.csv", package = "h2o")
australia = h2o.importFile(path = australia_path)
h2o.stopLogging()

## End(Not run)
```

---

**h2o.str**  
*Display the structure of an H2OFrame object*

**Description**

Display the structure of an H2OFrame object

**Usage**

```r
h2o.str(object, ..., cols = FALSE)
```

**Arguments**

- **object**: An H2OFrame.
- **...**: Further arguments to be passed from or to other methods.
- **cols**: Print the per-column str for the H2OFrame
Examples

```r
## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)
h2o.str(frame, cols = FALSE)

## End(Not run)
```

---

### h2o.stringdist

**Compute element-wise string distances between two H2OFrames**

#### Description

Compute element-wise string distances between two H2OFrames. Both frames need to have the same shape (N x M) and only contain string/factor columns. Return a matrix (H2OFrame) of shape N x M.

#### Usage

```r
h2o.stringdist(
  x,
  y,
  method = c("lv", "lcs", "qgram", "jaccard", "jw", "soundex"),
  compare_empty = TRUE
)
```

#### Arguments

- **x**: An H2OFrame
- **y**: A comparison H2OFrame
- **method**: A string identifier indicating what string distance measure to use. Must be one of: "lv" - Levenshtein distance "lcs" - Longest common substring distance "qgram" - q-gram distance "jaccard" - Jaccard distance between q-gram profiles "jw" - Jaro, or Jaro-Winker distance "soundex" - Distance based on soundex encoding
- **compare_empty**: if set to FALSE, empty strings will be handled as NaNs
Examples

```r
## Not run:
h2o.init()
x <- as.h2o(c("Martha", "Dwayne", "Dixon"))
y <- as.character(as.h2o(c("Marhta", "Duane", "Dicksonx")))
h2o.stringdist(x, y, method = "jw")

## End(Not run)
```

---

### h2o.strsplit

String Split

**Usage**

```r
h2o.strsplit(x, split)
```

**Arguments**

- `x`  
The column whose strings must be split.
- `split`  
The pattern to split on.

**Value**

An H2OFrame where each column is the outcome of the string split.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
string_to_split <- as.h2o("Split at every character.")
split_string <- h2o.strsplit(string_to_split, ")

## End(Not run)
```
**h2o.sub**

*String Substitute*

**Description**

Creates a copy of the target column in which each string has the first occurrence of the regex pattern replaced with the replacement substring.

**Usage**

```r
h2o.sub(pattern, replacement, x, ignore.case = FALSE)
```

**Arguments**

- `pattern`: The pattern to replace.
- `replacement`: The replacement pattern.
- `x`: The column on which to operate.
- `ignore.case`: Case sensitive or not

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
string_to_sub <- as.h2o("r tutorial")
sub_string <- h2o.sub("r ", "H2O ", string_to_sub)
## End(Not run)
```

---

**h2o.substring**

*Substring*

**Description**

Returns a copy of the target column that is a substring at the specified start and stop indices, inclusive. If the stop index is not specified, then the substring extends to the end of the original string. If start is longer than the number of characters in the original string, or is greater than stop, an empty string is returned. Negative start is coerced to 0.

**Usage**

```r
h2o.substring(x, start, stop = "[]")
h2o.substr(x, start, stop = "[]")
```
Arguments

- **x**: The column on which to operate.
- **start**: The index of the first element to be included in the substring.
- **stop**: Optional, The index of the last element to be included in the substring.

Examples

```r
# Not run:
library(h2o)
h2o.init()
string_to_substring <- as.h2o("1234567890")
substr <- h2o.substring(string_to_substring, 2) # Get substring from second index onwards
```

```
## End(Not run)
```

---

h2o.sum

*Compute the frame's sum by-column (or by-row).*

Description

Compute the frame’s sum by-column (or by-row).

Usage

```r
h2o.sum(x, na.rm = FALSE, axis = 0, return_frame = FALSE)
```

Arguments

- **x**: An H2OFrame object.
- **na.rm**: logical. indicating whether missing values should be removed.
- **axis**: An int that indicates whether to do down a column (0) or across a row (1). For row or column sums, the return_frame parameter must be TRUE.
- **return_frame**: A boolean that indicates whether to return an H2O frame or one single aggregated value. Default is FALSE.

See Also

- `sum` for the base R implementation.
Examples

```r
## Not run:
library(h2o)
#not run:

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)

h2o.sum(frame["C1"], na.rm = TRUE, axis = 0, return_frame = TRUE)
## End(Not run)
```

h2o.summary

---

**h2o.summary**

**Summarizes the columns of an H2OFrame.**

---

**Description**

A method for the `summary` generic. Summarizes the columns of an H2O data frame or subset of columns and rows using vector notation (e.g. dataset[row, col]).

**Usage**

```r
h2o.summary(object, factors = 6L, exact_quantiles = FALSE, ...)
```

## S3 method for class 'H2OFrame'
`summary(object, factors, exact_quantiles, ...)`

**Arguments**

- `object` An H2OFrame object.
- `factors` The number of factors to return in the summary. Default is the top 6.
- `exact_quantiles` Compute exact quantiles or use approximation. Default is to use approximation.
- `...` Further arguments passed to or from other methods.

**Details**

By default it uses approximated version of quantiles computation, however, user can modify this behavior by setting up `exact_quantiles` argument to true.

**Value**

A table displaying the minimum, 1st quartile, median, mean, 3rd quartile and maximum for each numeric column, and the levels and category counts of the levels in each categorical column.
Examples

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.importFile(path = prostate_path)
summary(prostate)
summary(prostate$GLEASON)
summary(prostate[, 4:6])
summary(prostate, exact_quantiles = TRUE)
## End(Not run)
```

h2o.svd

Singular value decomposition of an H2O data frame using the power method

Description

Singular value decomposition of an H2O data frame using the power method

Usage

```r
h2o.svd(
  training_frame,
  x,
  destination_key,
  model_id = NULL,
  validation_frame = NULL,
  ignore_const_cols = TRUE,
  score_each_iteration = FALSE,
  transform = c("NONE", "STANDARDIZE", "NORMALIZE", "DEMEAN", "DESCALE"),
  svd_method = c("GramSVD", "Power", "Randomized"),
  nv = 1,
  max_iterations = 1000,
  seed = -1,
  keep_u = TRUE,
  u_name = NULL,
  use_all_factor_levels = TRUE,
  max_runtime_secs = 0,
  export_checkpoints_dir = NULL
)
```

Arguments

- `training_frame`  Id of the training data frame.
- `x`              A vector containing the character names of the predictors in the model.
destination_key
(Optional) The unique key assigned to the resulting model. Automatically generated if none is provided.

model_id
Destination id for this model; auto-generated if not specified.

validation_frame
Id of the validation data frame.

ignore_const_cols
Logical. Ignore constant columns. Defaults to TRUE.

score_each_iteration
Logical. Whether to score during each iteration of model training. Defaults to FALSE.

transform
Transformation of training data Must be one of: "NONE", "STANDARDIZE", "NORMALIZE", "DEMEAN", "DESCALE". Defaults to NONE.

svd_method
Method for computing SVD (Caution: Randomized is currently experimental and unstable) Must be one of: "GramSVD", "Power", "Randomized". Defaults to GramSVD.

nv
Number of right singular vectors Defaults to 1.

max_iterations
Maximum iterations Defaults to 1000.

seed
Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).

keep_u
Logical. Save left singular vectors? Defaults to TRUE.

u_name
Frame key to save left singular vectors

use_all_factor_levels
Logical. Whether first factor level is included in each categorical expansion Defaults to TRUE.

max_runtime_secs
Maximum allowed runtime in seconds for model training. Use 0 to disable. Defaults to 0.

export_checkpoints_dir
Automatically export generated models to this directory.

Value

an object of class H2ODimReductionModel.

References

Examples

```r
## Not run:
library(h2o)
h2o.init()
australia_path <- system.file("extdata", "australia.csv", package = "h2o")
australia <- h2o.uploadFile(path = australia_path)
h2o.svd(training_frame = australia, nv = 8)

## End(Not run)
```

---

**h2o.table**  
Cross Tabulation and Table Creation in H2O

**Description**

Uses the cross-classifying factors to build a table of counts at each combination of factor levels.

**Usage**

```r
h2o.table(x, y = NULL, dense = TRUE)
```

**Arguments**

- `x`  
  An H2OFrame object with at most two columns.

- `y`  
  An H2OFrame similar to `x`, or `NULL`.

- `dense`  
  A logical for dense representation, which lists only non-zero counts, 1 combination per row. Set to `FALSE` to expand counts across all combinations.

**Value**

Returns a tabulated H2OFrame object.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
summary(prostate)
# Counts of the ages of all patients
head(h2o.table(prostate[, 3]))
h2o.table(prostate[, 3])

# Two-way table of ages (rows) and race (cols) of all patients
```
h2o.tabulate

Tabulation between Two Columns of an H2OFrame

Description
Simple Co-Occurrence based tabulation of X vs Y, where X and Y are two Vecs in a given dataset. Uses histogram of given resolution in X and Y. Handles numerical/categorical data and missing values. Supports observation weights.

Usage
h2o.tabulate(data, x, y, weights_column = NULL, nbins_x = 50, nbins_y = 50)

Arguments
- data: An H2OFrame object.
- x: predictor column
- y: response column
- weights_column: (optional) observation weights column
- nbins_x: number of bins for predictor column
- nbins_y: number of bins for response column

Value
Returns two TwoDimTables of 3 columns each count_table: X Y counts response_table: X meanY counts

Examples
## Not run:
library(h2o)
h2o.init()
df <- as.h2o(iris)
tab <- h2o.tabulate(data = df, x = "Sepal.Length", y = "Petal.Width",
weights_column = NULL, nbins_x = 10, nbins_y = 10)
plot(tab)

## End(Not run)
**h2o.tan**  
*Compute the tangent of x*

**Description**  
Compute the tangent of x

**Usage**  
```r
h2o.tan(x)
```

**Arguments**  
- `x`: An H2OFrame object.

**See Also**  
- `Trig` for the base R implementation, `tan()`.

**Examples**  
```r
## Not run:
library(h2o)
h2o.init()
frame <- h2o.createFrame(rows = 6, cols = 2,
                          categorical_fraction = 0.0,
                          missing_fraction = 0.7,
                          seed = 123)
h2o.tan(frame)
## End(Not run)
```

---

**h2o.tanh**  
*Compute the hyperbolic tangent of x*

**Description**  
Compute the hyperbolic tangent of x

**Usage**  
```r
h2o.tanh(x)
```

**Arguments**  
- `x`: An H2OFrame object.
See Also

Hyperbolic for the base R implementation, tanh().

Examples

```r
## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)
h2o.tanh(frame)
## End(Not run)
```

h2o.targetencoder

Transformation of a categorical variable with a mean value of the target variable

Description

Transformation of a categorical variable with a mean value of the target variable

Usage

```r
h2o.targetencoder(
  x,
  y,
  training_frame,
  model_id = NULL,
  fold_column = NULL,
  columns_to_encode = NULL,
  keep_original_categorical_columns = TRUE,
  blending = FALSE,
  inflection_point = 10,
  smoothing = 20,
  data_leakage_handling = c("leave_one_out", "k_fold", "none", "LeaveOneOut", "KFold", "None"),
  noise = 0.01,
  seed = -1,
  ...
)
```
h2o.targetencoder

Arguments

- **x** (Optional) A vector containing the names or indices of the predictor variables to use in building the model. If x is missing, then all columns except y are used.
- **y** The name or column index of the response variable in the data. The response must be either a numeric or a categorical/factor variable. If the response is numeric, then a regression model will be trained, otherwise it will train a classification model.
- **training_frame** Id of the training data frame.
- **model_id** Destination id for this model; auto-generated if not specified.
- **fold_column** Column with cross-validation fold index assignment per observation.
- **columns_to_encode** List of categorical columns or groups of categorical columns to encode. When groups of columns are specified, each group is encoded as a single column (interactions are created internally).
- **keep_original_categorical_columns** Logical. If true, the original non-encoded categorical features will remain in the result frame. Defaults to TRUE.
- **blending** Logical. If true, enables blending of posterior probabilities (computed for a given categorical value) with prior probabilities (computed on the entire set). This allows to mitigate the effect of categorical values with small cardinality. The blending effect can be tuned using the ‘inflection_point’ and ‘smoothing’ parameters. Defaults to FALSE.
- **inflection_point** Inflection point of the sigmoid used to blend probabilities (see ‘blending’ parameter). For a given categorical value, if it appears less than ‘inflection_point’ in a data sample, then the influence of the posterior probability will be smaller than the prior. Defaults to 10.
- **smoothing** Smoothing factor corresponds to the inverse of the slope at the inflection point on the sigmoid used to blend probabilities (see ‘blending’ parameter). If smoothing tends towards 0, then the sigmoid used for blending turns into a Heaviside step function. Defaults to 20.
- **data_leakage_handling** Data leakage handling strategy used to generate the encoding. Supported options are: 1) "none" (default) - no holdout, using the entire training frame. 2) "leave_one_out" - current row’s response value is subtracted from the per-level frequencies pre-calculated on the entire training frame. 3) "k_fold" - encodings for a fold are generated based on out-of-fold data. Must be one of: "leave_one_out", "k_fold", "none", "LeaveOneOut", "KFold", "None". Defaults to None.
- **noise** The amount of noise to add to the encoded column. Use 0 to disable noise, and -1 (=AUTO) to let the algorithm determine a reasonable amount of noise. Defaults to 0.01.
- **seed** Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).
- **...** Mainly used for backwards compatibility, to allow deprecated parameters.
### Examples

```r
## Not run:
library(h2o)
h2o.init()

# Import the titanic dataset
f <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/gbm_test/titanic.csv"
titanic <- h2o.importFile(f)

# Set response as a factor
response <- "survived"
titanic[response] <- as.factor(titanic[response])

# Split the dataset into train and test
splits <- h2o.splitFrame(data = titanic, ratios = .8, seed = 1234)
train <- splits[[1]]
test <- splits[[2]]

# Choose which columns to encode
encode_columns <- c("home.dest", "cabin", "embarked")

# Train a TE model
te_model <- h2o.targetencoder(x = encode_columns,
y = response,
training_frame = train,
fold_column = "pclass",
data_leakage_handling = "KFold")

# New target encoded train and test sets
train_te <- h2o.transform(te_model, train)
test_te <- h2o.transform(te_model, test)

## End(Not run)
```

---

**h2o.target_encode_apply**

*Apply Target Encoding Map to Frame*

**Description**


**Usage**

```r
h2o.target_encode_apply(  
data,  
x,
)```
```
y, 
target_encode_map, 
holdout_type, 
fold_column = NULL, 
blended_avg = TRUE, 
noise_level = NULL, 
seed = -1 
)

Arguments

data An H2OFrame object with which to apply the target encoding map.
x A list containing the names or indices of the variables to encode. A target encoding column will be created for each element in the list. Items in the list can be multiple columns. For example, if ‘x = list(c("A"), c("B", "C"))’, then the resulting frame will have a target encoding column for A and a target encoding column for B & C (in this case, we group by two columns).
y The name or column index of the response variable in the data. The response variable can be either numeric or binary.
target_encode_map A list of H2OFrame objects that is the results of the `h2o.target_encode_create` function.
holdout_type The holdout type used. Must be one of: "LeaveOneOut", "KFold", "None".
fold_column (Optional) The name or column index of the fold column in the data. Defaults to NULL (no ‘fold_column’). Only required if ‘holdout_type’ = "KFold".
blended_avg Logical. (Optional) Whether to perform blended average.
noise_level (Optional) The amount of random noise added to the target encoding. This helps prevent overfitting. Defaults to 0.01 * range of y.
seed (Optional) A random seed used to generate draws from the uniform distribution for random noise. Defaults to -1.

Value

Returns an H2OFrame object containing the target encoding per record.

See Also

h2o.target_encode_create for creating the target encoding map

Examples

```r
## Not run:
library(h2o)
h2o.init()

# Get Target Encoding Frame on bank-additional-full data with numeric `y`
data <- h2o.importFile(
  path = "https://s3.amazonaws.com/h2o-public-test-data/smalldata/demos/bank-additional-full.csv")
```
splits <- h2o.splitFrame(data, seed = 1234)
train <- splits[[1]]
test <- splits[[2]]
mapping <- h2o.target_encode_create(data = train, x = list(c("job"), c("job", "marital")), y = "age")

# Apply mapping to the training dataset
train_encode <- h2o.target_encode_apply(data = train, x = list(c("job"), c("job", "marital")), y = "age", mapping, holdout_type = "LeaveOneOut")

# Apply mapping to a test dataset
test_encode <- h2o.target_encode_apply(data = test, x = list(c("job"), c("job", "marital")), y = "age", target_encode_map = mapping, holdout_type = "None")

## End(Not run)

### h2o.target_encode_create

#### Create Target Encoding Map

**Description**

Creates a target encoding map based on group-by columns (\('x'\)) and a numeric or binary target column (\('y'\)). Computing target encoding for high cardinality categorical columns can improve performance of supervised learning models. A Target Encoding tutorial is available here: https://github.com/h2oai/h2o-tutorials/blob/master/best-practices/categorical-predictors/target_encoding.md.

**Usage**

h2o.target_encode_create(data, x, y, fold_column = NULL)

**Arguments**

- **data**: An H2OFrame object with which to create the target encoding map.
- **x**: A list containing the names or indices of the variables to encode. A target encoding map will be created for each element in the list. Items in the list can be multiple columns. For example, if \(x = \text{list}(\text{"A"}, \text{\text{\text{\"B", \"C\")}})\), then there will be one mapping frame for A and one mapping frame for B & C (in this case, we group by two columns).
- **y**: The name or column index of the response variable in the data. The response variable can be either numeric or binary.
- **fold_column** (Optional): The name or column index of the fold column in the data. Defaults to NULL (no ‘fold_column’).

**Value**

Returns a list of H2OFrame objects containing the target encoding mapping for each column in ‘x’.
See Also

h2o.target_encode_apply for applying the target encoding mapping to a frame.

Examples

```r
## Not run:
library(h2o)
h2o.init()

# Get Target Encoding Map on bank-additional-full data with numeric response
data <- h2o.importFile(
  path = "https://s3.amazonaws.com/h2o-public-test-data/smalldata/demos/bank-additional-full.csv")
mapping_age <- h2o.target_encode_create(data = data, x = list(c("job"), c("job", "marital")),
  y = "age")
head(mapping_age)

# Get Target Encoding Map on bank-additional-full data with binary response
mapping_y <- h2o.target_encode_create(data = data, x = list(c("job"), c("job", "marital")),
  y = "y")
head(mapping_y)

## End(Not run)
```

---

### h2o.tf_idf

Computes TF-IDF values for each word in given documents.

**Description**

Computes TF-IDF values for each word in given documents.

**Usage**

```r
h2o.tf_idf(
  frame,  # documents or words frame for which TF-IDF values should be computed.
  document_id_col,  # index or name of a column containing document IDs.
  text_col,  # index or name of a column containing documents if 'preprocess = TRUE' or words if 'preprocess = FALSE'.
  preprocess = TRUE,
  case_sensitive = TRUE
)
```

**Arguments**

- `frame`: documents or words frame for which TF-IDF values should be computed.
- `document_id_col`: index or name of a column containing document IDs.
- `text_col`: index or name of a column containing documents if `preprocess = TRUE` or words if `preprocess = FALSE`.
preprocess      whether input text data should be pre-processed. Defaults to ‘TRUE’.
case_sensitive whether input data should be treated as case sensitive. Defaults to ‘TRUE’.

Value
resulting frame with TF-IDF values. Row format: documentID, word, TF, IDF, TF-IDF

h2o.thresholds_and_metric_scores

Retrieve the thresholds and metric scores table

Description
Retrieves the thresholds and metric scores table from an H2OBinomialUpliftMetrics. The table contains indices, thresholds, all cumulative uplift values and cumulative number of observations. If “train” and “valid” parameters are FALSE (default), then the training table is returned. If more than one parameter is set to TRUE, then a named vector of tables is returned, where the names are “train”, “valid”.

Usage
h2o.thresholds_and_metric_scores(object, train = FALSE, valid = FALSE)

Arguments
object       An H2OBinomialUpliftMetrics
train        Retrieve the training thresholds and metric scores table
valid        Retrieve the validation thresholds and metric scores table

Examples
## Not run:
library(h2o)
h2o.init()
f <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/uplift/criteo_uplift_13k.csv"
train <- h2o.importFile(f)
train$treatment <- as.factor(train$treatment)
train$conversion <- as.factor(train$conversion)
model <- h2o.upliftRandomForest(training_frame=train, x=sprintf("f%s",seq(0:10)), y="conversion",
                                 ntree=10, max_depth=5, treatment_column="treatment",
                                 auuc_type="AUTO")
perf <- h2o.performance(model, train=TRUE)
h2o.thresholds_and_metric_scores(perf)
## End(Not run)
h2o.toFrame

Convert a word2vec model into an H2OFrame

Description

Converts a given word2vec model into an H2OFrame. The frame represents learned word embeddings.

Usage

h2o.toFrame(word2vec)

Arguments

word2vec A word2vec model.

Examples

```r
## Not run:
# Build a dummy word2vec model
data <- as.character(as.h2o(c("a", "b", "a")))
w2v_model <- h2o.word2vec(data, sent_sample_rate = 0, min_word_freq = 0, epochs = 1, vec_size = 2)

# Transform words to vectors and return average vector for each sentence
h2o.toFrame(w2v_model) # -> Frame made of 2 rows and 2 columns
## End(Not run)
```

h2o.tokenize

Tokenize String

Description

h2o.tokenize is similar to h2o.strsplit, the difference between them is that h2o.tokenize will store the tokenized text into a single column making it easier for additional processing (filtering stop words, word2vec algo, ...).

Usage

h2o.tokenize(x, split)

Arguments

x The column or columns whose strings to tokenize.

split The regular expression to split on.
**Value**

An H2OFrame with a single column representing the tokenized Strings. Original rows of the input DF are separated by NA.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
string_to_tokenize <- as.h2o("Split at every character and tokenize.")
tokenize_string <- h2o.tokenize(as.character(string_to_tokenize), ":")

## End(Not run)
```

---

**h2o.tolower**  
*Convert strings to lowercase*

**Description**

Convert strings to lowercase

**Usage**

`h2o.tolower(x)`

**Arguments**

*x*  
An H2OFrame object whose strings should be lower cased

**Value**

An H2OFrame with all entries in lowercase format

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
string_to_lower <- as.h2o("ABCDE")
lowered_string <- h2o.tolower(string_to_lower)

## End(Not run)
```
**h2o.topBottomN**

**H2O topBottomN**

**Description**

topBottomN function will grab the top N percent or bottom N percent of values of a column and return it in a H2OFrame.

**Usage**

```
h2o.topBottomN(x, column, nPercent, grabTopN)
```

**Arguments**

- **x**: an H2OFrame
- **column**: is a column name or column index to grab the top N percent value from
- **nPercent**: a top percentage values to grab
- **grabTopN**: if -1 grab bottom percentage, 1 grab top percentage

**Value**

An H2OFrame with 2 columns: first column is the original row indices, second column contains the values.

---

**h2o.topN**

**H2O topN**

**Description**

Extract the top N percent of values of a column and return it in a H2OFrame.

**Usage**

```
h2o.topN(x, column, nPercent)
```

**Arguments**

- **x**: an H2OFrame
- **column**: is a column name or column index to grab the top N percent value from
- **nPercent**: is a top percentage value to grab

**Value**

An H2OFrame with 2 columns. The first column is the original row indices, second column contains the topN values.
### Examples

```r
## Not run:
library(h2o)
h2o.init()

dataset <- h2o.importFile(f)
frameNames <- names(dataset)
nPercent <- c(1, 2, 3, 4)
nP <- nPercent[sample(1:length(nPercent), 1, replace = FALSE)]
colIndex <- sample(1:length(frameNames), 1, replace = FALSE)
h2o.topN(dataset, frameNames[colIndex], nP)

## End(Not run)
```

---

**h2o.totss**

Get the total sum of squares.

### Description

If "train", "valid", and "xval" parameters are FALSE (default), then the training totss value is returned. If more than one parameter is set to TRUE, then a named vector of totss' are returned, where the names are "train", "valid" or "xval".

### Usage

```r
h2o.totss(object, train = FALSE, valid = FALSE, xval = FALSE)
```

### Arguments

- **object**: An `H2OClusteringModel` object.
- **train**: Retrieve the training total sum of squares
- **valid**: Retrieve the validation total sum of squares
- **xval**: Retrieve the cross-validation total sum of squares

### Examples

```r
## Not run:
library(h2o)
h2o.init()

predictors <- c("sepal_len", "sepal_wid", "petal_len", "petal_wid")
km <- h2o.kmeans(x = predictors, training_frame = fr, k = 3, nfolds = 3)
h2o.totss(km, train = TRUE)

## End(Not run)
```
h2o.tot_withinss

Get the total within cluster sum of squares.

Description

If "train", "valid", and "xval" parameters are FALSE (default), then the training tot_withinss value is returned. If more than one parameter is set to TRUE, then a named vector of tot_withinss’ are returned, where the names are "train", "valid" or "xval".

Usage

h2o.tot_withinss(object, train = FALSE, valid = FALSE, xval = FALSE)

Arguments

object An H2OClusteringModel object.
train Retrieve the training total within cluster sum of squares
valid Retrieve the validation total within cluster sum of squares
xval Retrieve the cross-validation total within cluster sum of squares

Examples

## Not run:
library(h2o)
h2o.init()

predictors <- c("sepal_len", "sepal_wid", "petal_len", "petal_wid")
km <- h2o.kmeans(x = predictors, training_frame = fr, k = 3, nfolds = 3)
h2o.tot_withinss(km, train = TRUE)

## End(Not run)

h2o.toupper

Convert strings to uppercase

Description

Convert strings to uppercase

Usage

h2o.toupper(x)
h2o.train_segments

Arguments

- x: An H2OFrame object whose strings should be upper cased

Value

An H2OFrame with all entries in uppercase format

Examples

```r
## Not run:
library(h2o)
h2o.init()
string_to_upper <- as.h2o("abcde")
upper_string <- h2o.toupper(string_to_upper)

## End(Not run)
```

h2o.train_segments  

**H2O Segmented-Data Bulk Model Training**

Description

Provides a set of functions to train a group of models on different segments (subpopulations) of the training set.

Usage

```r
h2o.train_segments(
  algorithm,
  segment_columns,
  segment_models_id,
  parallelism = 1,
  ...
)
```

Arguments

- **algorithm**: Name of algorithm to use in training segment models (gbm, randomForest, kmeans, glm, deeplearning, naivebayes, psvm, xgboost, pca, svd, targetencoder, aggregator, word2vec, coxph, isolationforest, kmeans, stackedensemble, glrm, gam, anovaglm, modelselection).
- **segment_columns**: A list of columns to segment-by. H2O will group the training (and validation) dataset by the segment-by columns and train a separate model for each segment (group of rows).
- **segment_models_id**: Identifier for the returned collection of Segment Models. If not specified it will be automatically generated.
**parallelism**  Level of parallelism of bulk model building, it is the maximum number of models each H2O node will be building in parallel, defaults to 1.

...  Use to pass along training_frame parameter, x, y, and all non-default parameter values to the algorithm. Look at the specific algorithm - h2o.gbm, h2o.glm, h2o.kmeans, h2o.deepLearning - for available parameters.

**Details**  
Start Segmented-Data bulk Model Training for a given algorithm and parameters.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
iris_hf <- as.h2o(iris)
models <- h2o.train_segments(algorithm = "gbm",
    segment_columns = "Species",
    x = c(1:3), y = 4,
    training_frame = iris_hf,
    ntrees = 5,
    max_depth = 4)
as.data.frame(models)
## End(Not run)
```

---

**h2o.transform**  
*Use H2O Transformation model and apply the underlying transformation*

**Description**  
Use H2O Transformation model and apply the underlying transformation

**Usage**

```r
h2o.transform(model, ...)
```

**Arguments**

- **model**  
  A trained model representing the transformation strategy

- **...**  
  Transformation model-specific parameters

**Value**  
Returns an H2OFrame object with data transformed.
Applies target encoding to a given dataset

Usage

```r
## S4 method for signature 'H2OTargetEncoderModel'
h2o.transform(
  model,
  data,
  blending = NULL,
  inflection_point = -1,
  smoothing = -1,
  noise = NULL,
  as_training = FALSE,
  ...
)
```

Arguments

- **model**: A trained model representing the transformation strategy
- **data**: An H2OFrame with data to be transformed
- **blending**: Use blending during the transformation. Respects model settings when not set.
- **inflection_point**: Blending parameter. Only effective when blending is enabled. By default, model settings are respected, if not overridden by this setting.
- **smoothing**: Blending parameter. Only effective when blending is enabled. By default, model settings are respected, if not overridden by this setting.
- **noise**: An amount of random noise added to the encoding, this helps prevent overfitting. By default, model settings are respected, if not overridden by this setting.
- **as_training**: Must be set to True when encoding the training frame. Defaults to False.
- **...**: Mainly used for backwards compatibility, to allow deprecated parameters.

Value

Returns an H2OFrame object with data transformed.
Transform words (or sequences of words) to vectors using a word2vec model.

## S4 method for signature 'H2OWordEmbeddingModel'

```r
h2o.transform(model, words, aggregate_method = c("NONE", "AVERAGE"))
```

**Arguments**

- `model`: A word2vec model.
- `words`: An H2OFrame made of a single column containing source words.
- `aggregate_method`: Specifies how to aggregate sequences of words. If method is 'NONE' then no aggregation is performed and each input word is mapped to a single word-vector. If method is 'AVERAGE' then input is treated as sequences of words delimited by NA. Each word of a sequences is internally mapped to a vector and vectors belonging to the same sentence are averaged and returned in the result.

**Examples**

```r
# Not run:
h2o.init()

# Build a simple word2vec model
data <- as.character(as.h2o(c("a", "b", "a")))
w2v_model <- h2o.word2vec(data, sent_sample_rate = 0, min_word_freq = 0, epochs = 1, vec_size = 2)

# Transform words to vectors without aggregation
sentences <- as.character(as.h2o(c("b", "c", "a", NA, "b")))
h2o.transform(w2v_model, sentences) # -> 5 rows total, 2 rows NA ("c" is not in the vocabulary)

# Transform words to vectors and return average vector for each sentence
h2o.transform(w2v_model, sentences, aggregate_method = "AVERAGE") # -> 2 rows

# End(Not run)
```
h2o.transform_word2vec

Transform words (or sequences of words) to vectors using a word2vec model.

Description

Transform words (or sequences of words) to vectors using a word2vec model.

Usage

h2o.transform_word2vec(
  word2vec,
  words,
  aggregate_method = c("NONE", "AVERAGE")
)

Arguments

word2vec A word2vec model.
words An H2OFrame made of a single column containing source words.
aggregate_method Specifies how to aggregate sequences of words. If method is ‘NONE’ then no aggregation is performed and each input word is mapped to a single word-vector. If method is ‘AVERAGE’ then input is treated as sequences of words delimited by NA. Each word of a sequences is internally mapped to a vector and vectors belonging to the same sentence are averaged and returned in the result.

Examples

## Not run:
h2o.init()

# Build a dummy word2vec model
data <- as.character(as.h2o(c("a", "b", "a")))
w2v_model <- h2o.word2vec(data, sent_sample_rate = 0, min_word_freq = 0, epochs = 1, vec_size = 2)

# Transform words to vectors without aggregation
sentences <- as.character(as.h2o(c("b", "c", "a", NA, "b")))
h2o.transform(w2v_model, sentences) # -> 5 rows total, 2 rows NA ("c" is not in the vocabulary)

# Transform words to vectors and return average vector for each sentence
h2o.transform(w2v_model, sentences, aggregate_method = "AVERAGE") # -> 2 rows

## End(Not run)
h2o.trim

Trim Space

Description
Trim Space

Usage
h2o.trim(x)

Arguments
x The column whose strings should be trimmed.

Examples
## Not run:
library(h2o)
h2o.init()
string_to_trim <- as.h2o("r tutorial")
trim_string <- h2o.trim(string_to_trim)
## End(Not run)

h2o.trunc

Truncate values in x toward 0

Description
trunc takes a single numeric argument x and returns a numeric vector containing the integers formed by truncating the values in x toward 0.

Usage
h2o.trunc(x)

Arguments
x An H2OFrame object.

See Also
Round for the base R implementation, trunc().
Examples

```r
## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)
h2o.trunc(frame["C1"])

## End(Not run)
```

---

### h2o.unique

**H2O Unique**

#### Description

Extract unique values in the column.

#### Usage

```r
h2o.unique(x, include_nas = FALSE)
```

#### Arguments

- `x`: An H2OFrame object.
- `include_nas`: If set to TRUE, NAs are included. FALSE (turned off) by default.

#### Value

Returns an H2OFrame object.

#### Examples

```r
## Not run:
library(h2o)
h2o.init()

iris <- h2o.importFile(f)
h2o.unique(iris["class"])

## End(Not run)
```
h2o.upliftRandomForest

Build a Uplift Random Forest model

Description

Builds a Uplift Random Forest model on an H2OFrame.

Usage

```r
h2o.upliftRandomForest(
  x,
  y,
  training_frame,
  treatment_column,
  model_id = NULL,
  validation_frame = NULL,
  score_each_iteration = FALSE,
  score_tree_interval = 0,
  ignore_const_cols = TRUE,
  ntrees = 50,
  max_depth = 20,
  min_rows = 1,
  nbins = 20,
  nbins_top_level = 1024,
  nbins_cats = 1024,
  max_runtime_secs = 0,
  seed = -1,
  mtries = -2,
  sample_rate = 0.632,
  sample_rate_per_class = NULL,
  col_sample_rate_change_per_level = 1,
  col_sample_rate_per_tree = 1,
  histogram_type = c("AUTO", "UniformAdaptive", "Random", "QuantilesGlobal",
                    "RoundRobin"),
  categorical_encoding = c("AUTO", "Enum", "OneHotInternal", "OneHotExplicit",
                          "Binary", "Eigen", "LabelEncoder", "SortByResponse", "EnumLimited"),
  distribution = c("AUTO", "bernoulli", "multinomial", "gaussian", "poisson", "gamma",
                   "tweedie", "laplace", "quantile", "huber"),
  check_constant_response = TRUE,
  uplift_metric = c("AUTO", "KL", "Euclidean", "ChiSquared"),
  auuc_type = c("AUTO", "qini", "lift", "gain"),
  auuc_nbins = -1,
  verbose = FALSE
)
```
Arguments

x (Optional) A vector containing the names or indices of the predictor variables to use in building the model. If x is missing, then all columns except y are used.

y The name or column index of the response variable in the data. The response must be either a numeric or a categorical/factor variable. If the response is numeric, then a regression model will be trained, otherwise it will train a classification model.

training_frame Id of the training data frame.

treatment_column Define the column which will be used for computing uplift gain to select best split for a tree. The column has to divide the dataset into treatment (value 1) and control (value 0) groups. Defaults to treatment.

model_id Destination id for this model; auto-generated if not specified.

validation_frame Id of the validation data frame.

score_each_iteration Logical. Whether to score during each iteration of model training. Defaults to FALSE.

score_tree_interval Score the model after every so many trees. Disabled if set to 0. Defaults to 0.

ignore_const_cols Logical. Ignore constant columns. Defaults to TRUE.

ntrees Number of trees. Defaults to 50.

max_depth Maximum tree depth (0 for unlimited). Defaults to 20.

min_rows Fewest allowed (weighted) observations in a leaf. Defaults to 1.

nbins For numerical columns (real/int), build a histogram of (at least) this many bins, then split at the best point. Defaults to 20.

nbins_top_level For numerical columns (real/int), build a histogram of (at most) this many bins at the root level, then decrease by factor of two per level. Defaults to 1024.

nbins_cats For categorical columns (factors), build a histogram of this many bins, then split at the best point. Higher values can lead to more overfitting. Defaults to 1024.

max_runtime_secs Maximum allowed runtime in seconds for model training. Use 0 to disable. Defaults to 0.

seed Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).

mtries Number of variables randomly sampled as candidates at each split. If set to -1, defaults to $\sqrt{p}$ for classification and $p/3$ for regression (where $p$ is the # of predictors). Defaults to -2.

sample_rate Row sample rate per tree (from 0.0 to 1.0). Defaults to 0.632.
sample_rate_per_class
A list of row sample rates per class (relative fraction for each class, from 0.0 to 1.0), for each tree

col_sample_rate_change_per_level
Relative change of the column sampling rate for every level (must be > 0.0 and <= 2.0) Defaults to 1.

col_sample_rate_per_tree
Column sample rate per tree (from 0.0 to 1.0) Defaults to 1.

histogram_type
What type of histogram to use for finding optimal split points Must be one of: "AUTO", "UniformAdaptive", "Random", "QuantilesGlobal", "RoundRobin". Defaults to AUTO.

categorical_encoding
Encoding scheme for categorical features Must be one of: "AUTO", "Enum", "OneHotInternal", "OneHotExplicit", "Binary", "Eigen", "LabelEncoder", "Sort-ByResponse", "EnumLimited". Defaults to AUTO.

distribution
Distribution function Must be one of: "AUTO", "bernoulli", "multinomial", "gaussian", "poisson", "gamma", "tweedie", "laplace", "quantile", "huber". Defaults to AUTO.

check_constant_response
Logical. Check if response column is constant. If enabled, then an exception is thrown if the response column is a constant value. If disabled, then model will train regardless of the response column being a constant value or not. Defaults to TRUE.

uplift_metric
Divergence metric used to find best split when building an uplift tree. Must be one of: "AUTO", "KL", "Euclidean", "ChiSquared". Defaults to AUTO.

auuc_type
Metric used to calculate Area Under Uplift Curve. Must be one of: "AUTO", "qini", "lift", "gain". Defaults to AUTO.

auuc_nbins
Number of bins to calculate Area Under Uplift Curve. Defaults to -1.

verbose
Logical. Print scoring history to the console (Metrics per tree). Defaults to FALSE.

Value
Creates a H2OModel object of the right type.

See Also
predict.H2OModel for prediction
**h2o.upload_model**

Upload a binary model from the provided local path to the H2O cluster. (H2O model can be saved in a binary form either by `saveModel()` or by `download_model()` function.)

**Description**

Upload a binary model from the provided local path to the H2O cluster. (H2O model can be saved in a binary form either by `saveModel()` or by `download_model()` function.)

**Usage**

```r
h2o.upload_model(path)
```

**Arguments**

- `path` A path on the machine this python session is currently connected to, specifying the location of the model to upload.

**Value**

Returns a new `H2OModel` object.

**See Also**

- `h2o.saveModel`, `h2o.download_model`

---

**h2o.upload_mojo**

Imports a MOJO from a local filesystem, creating a Generic model with it.

**Description**

Usage example: `mojo_model <- h2o.upload_mojo(model_file_path = "/path/to/local/mojo.zip")
predictions <- h2o.predict(mojo_model, dataset)`

**Usage**

```r
h2o.upload_mojo(mojo_local_file_path, model_id = NULL)
```

**Arguments**

- `mojo_local_file_path` Filesystem path to the model imported
- `model_id` Model ID, default is NULL.
h2o.var

Variance of a column or covariance of columns.

Description

Compute the variance or covariance matrix of one or two H2OFrames.

Usage

h2o.var(x, y = NULL, na.rm = FALSE, use)

var(x, y = NULL, na.rm = FALSE, use)

Arguments

x

An H2OFrame object.

y

NULL (default) or an H2OFrame. The default is equivalent to y = x.

na.rm

logical. Should missing values be removed?

use

An optional character string indicating how to handle missing values. This must be one of the following: "everything" - outputs NaNs whenever one of its contributing observations is missing "all.obs" - presence of missing observations will throw an error "complete.obs" - discards missing values along with all observations in their rows so that only complete observations are used.
h2o.varimp

See Also
cor for the base R implementation, var(). h2o.sd for standard deviation.

Examples

## Not run:
library(h2o)
h2o.init()

prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
var(prostate$AGE)

## End(Not run)

---

h2o.varimp Retrieve the variable importance.

Description
Retrieve the variable importance.

Usage

h2o.varimp(object, ...)

Arguments

object An H2O object.

... Additional arguments for specific use-cases.

Examples

## Not run:
library(h2o)
h2o.init()

f <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/prostate/prostate_complete.csv.zip"
pros <- h2o.importFile(f)
response <- "GLEASON"
predictors <- c("ID", "AGE", "CAPSULE", "DCAPS", "PSA", "VOL", "DPROS")
aml <- h2o.automl(x = predictors, y = response, training_frame = pros, max_runtime_secs = 60)

h2o.varimp(aml, top_n = 20) # get variable importance matrix for the top 20 models

h2o.varimp(aml@leader) # get variable importance for the leader model

## End(Not run)
h2o.varimp,H2OAutoML-method

Retrieve the variable importance.

Description

Retrieve the variable importance.

Usage

## S4 method for signature 'H2OAutoML'

h2o.varimp(object, top_n = 20)

Arguments

- **object**: An H2OAutoML object.
- **top_n**: Show at most top_n models.

Examples

## Not run:
library(h2o)
h2o.init()
pros <- h2o.importFile(f)
response <- "GLEASON"
predictors <- c("ID", "AGE", "CAPSULE", "DCAPS", "PSA", "VOL", "DPROS")
aml <- h2o.automl(x = predictors, y = response, training_frame = pros, max_runtime_secs = 60)
h2o.varimp(aml)

## End(Not run)

h2o.varimp,H2OFrame-method

Retrieve the variable importance.

Description

Retrieve the variable importance.

Usage

## S4 method for signature 'H2OFrame'

h2o.varimp(object)
Arguments

object A leaderboard frame.

Examples

## Not run:
library(h2o)
h2o.init()

f <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/prostate/prostate_complete.csv.zip"
pros <- h2o.importFile(f)
response <- "GLEASON"
predictors <- c("ID", "AGE", "CAPSULE", "DCAPS", "PSA", "VOL", "DPROS")
aml <- h2o.automl(x = predictors, y = response, training_frame = pros, max_runtime_secs = 60)
h2o.varimp(aml@leaderboard[1:5,])

## End(Not run)

h2o.varimp,H2OModel-method

Retrieve the variable importance.

Description

Retrieve the variable importance.

Usage

## S4 method for signature 'H2OModel'
h2o.varimp(object)

Arguments

object An H2OModel object.

Examples

## Not run:
library(h2o)
h2o.init()

f <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/prostate/prostate_complete.csv.zip"
pros <- h2o.importFile(f)
response <- "GLEASON"
predictors <- c("ID", "AGE", "CAPSULE", "DCAPS", "PSA", "VOL", "DPROS")
model <- h2o.glm(x = predictors, y = response, training_frame = pros)
h2o.varimp(model)

## End(Not run)
Variable Importance Heatmap across multiple models

Variable importance heatmap shows variable importance across multiple models. Some models in H2O return variable importance for one-hot (binary indicator) encoded versions of categorical columns (e.g. Deep Learning, XGBoost). In order for the variable importance of categorical columns to be compared across all model types we compute a summarization of the the variable importance across all one-hot encoded features and return a single variable importance for the original categorical feature. By default, the models and variables are ordered by their similarity.

Usage

h2o.varimp_heatmap(object, top_n = 20)

Arguments

- object: A list of H2O models, an H2O AutoML instance, or an H2OFrame with a 'model_id' column (e.g. H2OAutoML leaderboard).
- top_n: Integer specifying the number models shown in the heatmap (based on leaderboard ranking). Defaults to 20.

Value

A ggplot2 object.

Examples

```r
## Not run:
library(h2o)
h2o.init()

# Import the wine dataset into H2O:
df <- h2o.importFile(f)

# Set the response
response <- "quality"

# Split the dataset into a train and test set:
splits <- h2o.splitFrame(df, ratios = 0.8, seed = 1)
train <- splits[[1]]
test <- splits[[2]]

# Build and train the model:
aml <- h2o.automl(y = response,
                 training_frame = train,
                 max_models = 10,
                 top_n = 20)
```

# Create the variable importance heatmap

```r
varimp_heatmap <- h2o.varimp_heatmap(aml)
print(varimp_heatmap)
```

## End(Not run)

---

### h2o.varimp_plot

**Plot Variable Importances**

**Description**

Plot Variable Importances

**Usage**

```r
h2o.varimp_plot(model, num_of_features = NULL)
```

**Arguments**

- `model`: A trained model (accepts a trained random forest, GBM, or deep learning model, will use `h2o.std_coef_plot` for a trained GLM
- `num_of_features`: The number of features shown in the plot (default is 10 or all if less than 10).

**See Also**

- `h2o.std_coef_plot` for GLM.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.importFile(prostate_path)
prostate[, 2] <- as.factor(prostate[, 2])
model <- h2o.gbm(x = 3:9, y = 2, training_frame = prostate, distribution = "bernoulli")
h2o.varimp_plot(model)

# for deep learning set the variable_importance parameter to TRUE
iris_hf <- as.h2o(iris)
iris_dl <- h2o.deeplearning(x = 1:4, y = 5, training_frame = iris_hf,
variable_importances = TRUE)
h2o.varimp_plot(iris_dl)

## End(Not run)
```
h2o.varsplits

Retrieve per-variable split information for a given Isolation Forest model. Output will include:
- count - The number of times a variable was used to make a split.
- aggregated_split_ratios - The split ratio is defined as \(\frac{\text{abs}(\#\text{left}_\text{observations} - \#\text{right}_\text{observations})}{\#\text{before}_\text{split}}\). Even splits (\#\text{left}_\text{observations} \approx \#\text{right}_\text{observations}) contribute less to the total aggregated split ratio value for the given feature; highly imbalanced splits (eg. \#\text{left}_\text{observations} \gg \#\text{right}_\text{observations}) contribute more.
- aggregated_split_depths - The sum of all depths of a variable used to make a split. (If a variable is used on level \(N\) of a tree, then it contributes with \(N\) to the total aggregate.)

Usage

h2o.varsplits(object)

Arguments

object

An Isolation Forest model represented by H2OModel object.

h2o.week

Convert Milliseconds to Week of Week Year in H2O Datasets

Description

Converts the entries of an H2OFrame object from milliseconds to weeks of the week year (starting from 1).
**h2o.week**

**Usage**

```r
h2o.week(x)
```

**Arguments**

- `x` An H2OFrame object.

**Value**

An H2OFrame object containing the entries of `x` converted to weeks of the week year.

**See Also**

- `h2o.month`

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

hdf <- h2o.importFile(f)
h2o.week(hdf["ds9"])
## End(Not run)
```

---

**h2o.weights**

*Retrieve the respective weight matrix*

**Description**

Retrieve the respective weight matrix

**Usage**

```r
h2o.weights(object, matrix_id = 1)
```

**Arguments**

- `object` An H2OModel or H2OModelMetrics
- `matrix_id` An integer, ranging from 1 to number of layers + 1, that specifies the weight matrix to return.
**Examples**

```r
## Not run:
library(h2o)
h2o.init()

census <- h2o.importFile(f)
census[, 1] <- as.factor(census[, 1])
dl_model <- h2o.deeplearning(x = c(1:3), y = 4, training_frame = census,
                              hidden = c(17, 191),
                              epochs = 1,
                              balance_classes = FALSE,
                              export_weights_and_biases = TRUE)
h2o.weights(dl_model, matrix_id = 1)

## End(Not run)
```

**h2o.which**

*Which indices are TRUE?*

**Description**

Give the TRUE indices of a logical object, allowing for array indices.

**Usage**

```r
h2o.which(x)
```

**Arguments**

- `x` An H2OFrame object.

**Value**

Returns an H2OFrame object.

**See Also**

`which` for the base R method.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

iris_hf <- as.h2o(iris)
h2o.which(iris_hf[, 1] == 4.4)

## End(Not run)
```
Description

Get the index of the max value in a column or row

Usage

h2o.which_max(x, na.rm = TRUE, axis = 0)

which.max.H2OFrame(x, na.rm = TRUE, axis = 0)

which.min.H2OFrame(x, na.rm = TRUE, axis = 0)

Arguments

- **x**: An H2OFrame object.
- **na.rm**: logical. Indicate whether missing values should be removed.
- **axis**: integer. Indicate whether to calculate the mean down a column (0) or across a row (1).

Value

Returns an H2OFrame object.

See Also

- which.min for the base R method, which.max().

Examples

```r
## Not run:
library(h2o)
h2o.init()

census <- h2o.importFile(f)
census[, 1] <- as.factor(census[, 1])
dl_model <- h2o.deeplearning(x = c(1:3), y = 4, hidden = c(17, 191),
  epochs = 1, training_frame = census,
  balance_classes = FALSE,
  export_weights_and_biases = TRUE)
h2o.which_max(census["PER CAPITA INCOME"], na.rm = FALSE, axis = 0)
## End(Not run)
```
**h2o.which_min**

Which index contains the min value?

**Description**

Get the index of the min value in a column or row

**Usage**

`h2o.which_min(x, na.rm = TRUE, axis = 0)`

**Arguments**

- **x** An H2OFrame object.
- **na.rm** logical. Indicate whether missing values should be removed.
- **axis** integer. Indicate whether to calculate the mean down a column (0) or across a row (1).

**Value**

Returns an H2OFrame object.

**See Also**

[which.min](https://www.rdocumentation.org/packages/base/versions/3.6.2/topics/which.min) for the base R method.

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

census <- h2o.importFile(f)
dl_model <- h2o.deeplearning(x = c(1:3), y = 4, hidden = c(17, 191),
                             epochs = 1, training_frame = census,
                             balance_classes = FALSE,
                             export_weights_and_biases = TRUE)
h2o.which_min(census["PER CAPITA INCOME "], na.rm = FALSE, axis = 0)

## End(Not run)
```
**h2o.withinss**

Get the Within SS

**Description**

Get the Within SS

**Usage**

```
h2o.withinss(object)
```

**Arguments**

- `object`: An `H2OClusteringModel` object.

---

**h2o.word2vec**

Trains a word2vec model on a String column of an H2O data frame

**Description**

Trains a word2vec model on a String column of an H2O data frame

**Usage**

```
h2o.word2vec(
    training_frame = NULL,
    model_id = NULL,
    min_word_freq = 5,
    word_model = c("SkipGram", "CBOW"),
    norm_model = c("HSM"),
    vec_size = 100,
    window_size = 5,
    sent_sample_rate = 0.001,
    init_learning_rate = 0.025,
    epochs = 5,
    pre_trained = NULL,
    max_runtime_secs = 0,
    export_checkpoints_dir = NULL
)
```
Arguments

**training_frame**  
Id of the training data frame.

**model_id**  
Destination id for this model; auto-generated if not specified.

**min_word_freq**  
This will discard words that appear less than <int> times. Defaults to 5.

**word_model**  
The word model to use (SkipGram or CBOW). Must be one of: "SkipGram", "CBOW". Defaults to SkipGram.

**norm_model**  
Use Hierarchical Softmax. Must be one of: "HSM". Defaults to HSM.

**vec_size**  
Set size of word vectors. Defaults to 100.

**window_size**  
Set max skip length between words. Defaults to 5.

**sent_sample_rate**  
Set threshold for occurrence of words. Those that appear with higher frequency in the training data will be randomly down-sampled; useful range is (0, 1e-5). Defaults to 0.001.

**init_learning_rate**  
Set the starting learning rate. Defaults to 0.025.

**epochs**  
Number of training iterations to run. Defaults to 5.

**pre_trained**  
Id of a data frame that contains a pre-trained (external) word2vec model.

**max_runtime_secs**  
Maximum allowed runtime in seconds for model training. Use 0 to disable. Defaults to 0.

**export_checkpoints_dir**  
Automatically export generated models to this directory.

Examples

```r
## Not run:
library(h2o)
h2o.init()

# Import the CraigslistJobTitles dataset
                            col.names = c("category", "jobtitle"), col.types = c("String", "String"), header = TRUE)

# Build and train the Word2Vec model
words <- h2o.tokenize(job_titles, " ")
vec <- h2o.word2vec(training_frame = words)
h2o.findSynonyms(vec, "teacher", count = 20)

## End(Not run)
```
Build an eXtreme Gradient Boosting model

Description

Builds a eXtreme Gradient Boosting model using the native XGBoost backend.

Usage

```r
h2o.xgboost(
  x,
  y,
  training_frame,
  model_id = NULL,
  validation_frame = NULL,
  nfolds = 0,
  keep_cross_validation_models = TRUE,
  keep_cross_validation_predictions = FALSE,
  keep_cross_validation_fold_assignment = FALSE,
  score_each_iteration = FALSE,
  fold_assignment = c("AUTO", "Random", "Modulo", "Stratified"),
  fold_column = NULL,
  ignore_const_cols = TRUE,
  offset_column = NULL,
  weights_column = NULL,
  stopping_rounds = 0,
  stopping_metric = c("AUTO", "deviance", "logloss", "MSE", "RMSE", "MAE", "RMSLE", "AUC", "AUCPR", "lift_top_group", "misclassification", "mean_per_class_error", "custom", "custom_increasing"),
  stopping_tolerance = 0.001,
  max_runtime_secs = 0,
  seed = -1,
  distribution = c("AUTO", "bernoulli", "multinomial", "gaussian", "poisson", "gamma", "tweedie", "laplace", "quantile", "huber"),
  tweedie_power = 1.5,
  quiet_mode = TRUE,
  checkpoint = NULL,
  export_checkpoints_dir = NULL,
  ntrees = 50,
  max_depth = 6,
  min_rows = 1,
  min_child_weight = 1,
  learn_rate = 0.3,
  eta = 0.3,
  sample_rate = 1,
  ...)```

subsample = 1,
col_sample_rate = 1,
colsample_bylevel = 1,
col_sample_rate_per_tree = 1,
colsample_bytree = 1,
colsample_bynode = 1,
max_abs_leafnode_pred = 0,
max_delta_step = 0,
monotone_constraints = NULL,
interaction_constraints = NULL,
score_tree_interval = 0,
min_split_improvement = 0,
gamma = 0,
nthread = -1,
save_matrix_directory = NULL,
build_tree_one_node = FALSE,
calibrate_model = FALSE,
calibration_frame = NULL,
max_bins = 256,
max_leaves = 0,
sample_type = c("uniform", "weighted"),
normalize_type = c("tree", "forest"),
rate_drop = 0,
one_drop = FALSE,
skip_drop = 0,
tree_method = c("auto", "exact", "approx", "hist"),
grow_policy = c("depthwise", "lossguide"),
booster = c("gbtree", "gblinear", "dart"),
reg_lambda = 1,
reg_alpha = 0,
dmatrix_type = c("auto", "dense", "sparse"),
backend = c("auto", "gpu", "cpu"),
gpu_id = NULL,
gainslift_bins = -1,
auc_type = c("AUTO", "NONE", "MACRO_OVR", "WEIGHTED_OVR", "MACRO_OVO",
           "WEIGHTED_OVO"),
scale_pos_weight = 1,
verbose = FALSE
)

Arguments

x (Optional) A vector containing the names or indices of the predictor variables to use in building the model. If x is missing, then all columns except y are used.

y The name or column index of the response variable in the data. The response must be either a numeric or a categorical/factor variable. If the response is numeric, then a regression model will be trained, otherwise it will train a classification model.
training_frame  Id of the training data frame.
model_id         Destination id for this model; auto-generated if not specified.
validation_frame Id of the validation data frame.
nfolds           Number of folds for K-fold cross-validation (0 to disable or >= 2). Defaults to 0.
keep_cross_validation_models Logical. Whether to keep the cross-validation models. Defaults to TRUE.
keep_cross_validation_predictions Logical. Whether to keep the predictions of the cross-validation models. Defaults to FALSE.
keep_cross_validation_fold_assignment Logical. Whether to keep the cross-validation fold assignment. Defaults to FALSE.
score_eachIteration Logical. Whether to score during each iteration of model training. Defaults to FALSE.
fold_assignment  Cross-validation fold assignment scheme, if fold_column is not specified. The 'Stratified' option will stratify the folds based on the response variable, for classification problems. Must be one of: "AUTO", "Random", "Modulo", "Stratified". Defaults to AUTO.
fold_column      Column with cross-validation fold index assignment per observation.
ignore_const_cols Logical. Ignore constant columns. Defaults to TRUE.
offset_column    Offset column. This will be added to the combination of columns before applying the link function.
weights_column   Column with observation weights. Giving some observation a weight of zero is equivalent to excluding it from the dataset; giving an observation a relative weight of 2 is equivalent to repeating that row twice. Negative weights are not allowed. Note: Weights are per-row observation weights and do not increase the size of the data frame. This is typically the number of times a row is repeated, but non-integer values are supported as well. During training, rows with higher weights matter more, due to the larger loss function pre-factor. If you set weight = 0 for a row, the returned prediction frame at that row is zero and this is incorrect. To get an accurate prediction, remove all rows with weight == 0.
stopping_rounds  Early stopping based on convergence of stopping_metric. Stop if simple moving average of length k of the stopping_metric does not improve for k:=stopping_rounds scoring events (0 to disable) Defaults to 0.
stopping_metric  Metric to use for early stopping (AUTO: logloss for classification, deviance for regression and anomaly_score for Isolation Forest). Note that custom and custom_increasing can only be used in GBM and DRF with the Python client. Must be one of: "AUTO", "deviance", "logloss", "MSE", "RMSE", "MAE", "RMSLE", "AUC", "AUCPR", "lift_top_group", "misclassification", "mean_per_class_error", "custom", "custom_increasing". Defaults to AUTO.
stopping_tolerance
    Relative tolerance for metric-based stopping criterion (stop if relative improvement is not at least this much) Defaults to 0.001.

max_runtime_secs
    Maximum allowed runtime in seconds for model training. Use 0 to disable. Defaults to 0.

seed
    Seed for random numbers (affects certain parts of the algo that are stochastic and those might or might not be enabled by default). Defaults to -1 (time-based random number).

distribution
    Distribution function Must be one of: "AUTO", "bernoulli", "multinomial", "gaussian", "poisson", "gamma", "tweedie", "laplace", "quantile", "huber". Defaults to AUTO.

tweedie_power
    Tweedie power for Tweedie regression, must be between 1 and 2. Defaults to 1.5.

categorical_encoding
    Encoding scheme for categorical features Must be one of: "AUTO", "Enum", "OneHotInternal", "OneHotExplicit", "Binary", "Eigen", "LabelEncoder", "Sort-ByResponse", "EnumLimited". Defaults to AUTO.

quiet_mode
    Logical. Enable quiet mode Defaults to TRUE.

checkpoint
    Model checkpoint to resume training with.

export_checkpoints_dir
    Automatically export generated models to this directory.

ntrees
    (same as n_estimators) Number of trees. Defaults to 50.

max_depth
    Maximum tree depth (0 for unlimited). Defaults to 6.

min_rows
    (same as min_child_weight) Fewest allowed (weighted) observations in a leaf. Defaults to 1.

min_child_weight
    (same as min_rows) Fewest allowed (weighted) observations in a leaf. Defaults to 1.

learn_rate
    (same as eta) Learning rate (from 0.0 to 1.0) Defaults to 0.3.

eta
    (same as learn_rate) Learning rate (from 0.0 to 1.0) Defaults to 0.3.

sample_rate
    (same as subsample) Row sample rate per tree (from 0.0 to 1.0) Defaults to 1.

subsample
    (same as sample_rate) Row sample rate per tree (from 0.0 to 1.0) Defaults to 1.

col_sample_rate
    (same as colsample_bylevel) Column sample rate (from 0.0 to 1.0) Defaults to 1.

colsample_bylevel
    (same as col_sample_rate) Column sample rate (from 0.0 to 1.0) Defaults to 1.

col_sample_rate_per_tree
    (same as colsample_bytree) Column sample rate per tree (from 0.0 to 1.0) Defaults to 1.

colsample_bytree
    (same as col_sample_rate_per_tree) Column sample rate per tree (from 0.0 to 1.0) Defaults to 1.
colsample_bynode
  Column sample rate per tree node (from 0.0 to 1.0) Defaults to 1.

max_abs_leafnode_pred
  (same as max_delta_step) Maximum absolute value of a leaf node prediction
  Defaults to 0.0.

max_delta_step
  (same as max_abs_leafnode_pred) Maximum absolute value of a leaf node pre-
  diction Defaults to 0.0.

monotone_constraints
  A mapping representing monotonic constraints. Use +1 to enforce an increasing
  constraint and -1 to specify a decreasing constraint.

interaction_constraints
  A set of allowed column interactions.

score_tree_interval
  Score the model after every so many trees. Disabled if set to 0. Defaults to 0.

min_split_improvement
  (same as gamma) Minimum relative improvement in squared error reduction for
  a split to happen Defaults to 0.0.

gamma
  (same as min_split_improvement) Minimum relative improvement in squared
  error reduction for a split to happen Defaults to 0.0.

nthread
  Number of parallel threads that can be used to run XGBoost. Cannot exceed
  H2O cluster limits (-nthreads parameter). Defaults to maximum available De-
  fault to -1.

save_matrix_directory
  Directory where to save matrices passed to XGBoost library. Useful for debug-
  ging.

build_tree_one_node
  Logical. Run on one node only; no network overhead but fewer cpus used.
  Suitable for small datasets. Defaults to FALSE.

calibrate_model
  Logical. Use Platt Scaling to calculate calibrated class probabilities. Cali-
  bration can provide more accurate estimates of class probabilities. Defaults to
  FALSE.

calibration_frame
  Calibration frame for Platt Scaling

max_bins
  For tree_method=hist only: maximum number of bins Defaults to 256.

max_leaves
  For tree_method=hist only: maximum number of leaves Defaults to 0.

sample_type
  For booster=dart only: sample_type Must be one of: "uniform", "weighted". Defaults to uniform.

normalize_type
  For booster=dart only: normalize_type Must be one of: "tree", "forest". Defaults to tree.

rate_drop
  For booster=dart only: rate_drop (0..1) Defaults to 0.0.

one_drop
  Logical. For booster=dart only: one_drop Defaults to FALSE.

skip_drop
  For booster=dart only: skip_drop (0..1) Defaults to 0.0.

tree_method
  Tree method Must be one of: "auto", "exact", "approx", "hist". Defaults to auto.
grow_policy  Grow policy - depthwise is standard GBM, lossguide is LightGBM. Must be one of: "depthwise", "lossguide". Defaults to depthwise.
booster  Booster type. Must be one of: "gbtree", "gblinear", "dart". Defaults to gbtree.
reg_lambda  L2 regularization. Defaults to 1.0.
reg_alpha  L1 regularization. Defaults to 0.0.
dmatrix_type  Type of DMatrix. For sparse, NAs and 0 are treated equally. Must be one of: "auto", "dense", "sparse". Defaults to auto.
backend  Backend. By default (auto), a GPU is used if available. Must be one of: "auto", "gpu", "cpu". Defaults to auto.
gpu_id  Which GPU(s) to use.
gainslift_bins  Gains/Lift table number of bins. 0 means disabled. Default value -1 means automatic binning. Defaults to -1.
auc_type  Set default multinomial AUC type. Must be one of: "AUTO", "NONE", "MACRO_OVR", "WEIGHTED_OVR", "MACRO_OVO", "WEIGHTED_OVO". Defaults to AUTO.
scale_pos_weight  Controls the effect of observations with positive labels in relation to the observations with negative labels on gradient calculation. Useful for imbalanced problems. Defaults to 1.0.
verbose  Logical. Print scoring history to the console (Metrics per tree). Defaults to FALSE.

Examples

```r
## Not run:
library(h2o)
h2o.init()

# Import the titanic dataset
f <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/gbm_test/titanic.csv"
titanic <- h2o.importFile(f)

# Set predictors and response; set response as a factor
titanic['survived'] <- as.factor(titanic['survived'])
predictors <- setdiff(colnames(titanic), colnames(titanic)[2:3])
response <- "survived"

# Split the dataset into train and valid
splits <- h2o.splitFrame(data = titanic, ratios = .8, seed = 1234)
train <- splits[[1]]
valid <- splits[[2]]

# Train the XGB model
titanic_xgb <- h2o.xgboost(x = predictors, y = response,
                          training_frame = train, validation_frame = valid,
                          booster = "dart", normalize_type = "tree",
                          seed = 1234)

## End(Not run)
```
**h2o.xgboost.available**  
*Determines whether an XGBoost model can be built*

**Description**
Ask the H2O server whether a XGBoost model can be built. (Depends on availability of native backend.) Returns True if a XGBoost model can be built, or False otherwise.

**Usage**
```
h2o.xgboost.available()
```

---

**h2o.year**  
*Convert Milliseconds to Years in H2O Datasets*

**Description**
Convert the entries of an H2OFrame object from milliseconds to years, indexed starting from 1900.

**Usage**
```
h2o.year(x)
year(x)
```

```
## S3 method for class 'H2OFrame'
year(x)
```

**Arguments**
- `x`  
  An H2OFrame object.

**Details**
This method calls the function of the MutableDateTime class in Java.

**Value**
An H2OFrame object containing the entries of `x` converted to years

**See Also**
- `h2o.month`
Examples

```r
## Not run:
library(h2o)
h2o.init()
hdf <- h2o.importFile(f)
h2o.year(hdf["ds9"])

## End(Not run)
```

H2OAutoML-class  

The **H2OAutoML** class

Description

This class represents an H2OAutoML object.

H2OClusteringModel-class  

The **H2OClusteringModel** object.

Description

This virtual class represents a clustering model built by H2O.

Details

This object has slots for the key, which is a character string that points to the model key existing in the H2O cluster, the data used to build the model (an object of class H2OFrame).

Slots

- `model_id` A character string specifying the key for the model fit in the H2O cluster’s key-value store.
- `algorithm` A character string specifying the algorithm that was used to fit the model.
- `parameters` A list containing the parameter settings that were used to fit the model that differ from the defaults.
- `allparameters` A list containing all parameters used to fit the model.
- `model` A list containing the characteristics of the model returned by the algorithm.
  - `size` The number of points in each cluster.
  - `totss` Total sum of squared error to grand mean.
  - `withinss` A vector of within-cluster sum of squared error.
  - `tot_withinss` Total within-cluster sum of squared error.
  - `betweenss` Between-cluster sum of squared error.
The H2OConnection class.

Description

This class represents a connection to an H2O cluster.

Usage

```r
## S4 method for signature 'H2OConnection'
show(object)
```

Arguments

- `object`: an H2OConnection object.

Details

Because H2O is not a master-slave architecture, there is no restriction on which H2O node is used to establish the connection between R (the client) and H2O (the server).

A new H2O connection is established via the `h2o.init()` function, which takes as parameters the `ip` and `port` of the machine running an instance to connect with. The default behavior is to connect with a local instance of H2O at port 54321, or to boot a new local instance if one is not found at port 54321.

Slots

- `ip`: A character string specifying the IP address of the H2O cluster.
- `port`: A numeric value specifying the port number of the H2O cluster.
- `name`: A character value specifying the name of the H2O cluster.
- `proxy`: A character specifying the proxy path of the H2O cluster.
- `https`: Set this to TRUE to use https instead of http.
- `cacert`: Path to a CA bundle file with root and intermediate certificates of trusted CAs.
- `insecure`: Set this to TRUE to disable SSL certificate checking.
- `username`: Username to login with.
- `password`: Password to login with.
- `use_spnego`: Set this to TRUE to use SPNEGO authentication.
- `cookies`: Cookies to add to request
- `context_path`: Context path which is appended to H2O server location.
- `mutable`: An H2OConnectionMutableState object to hold the mutable state for the H2O connection.
H2OConnectionMutableState

The H2OConnectionMutableState class

Description
This class represents the mutable aspects of a connection to an H2O cluster.

Slots

session_id  A character string specifying the H2O session identifier.
key_count  A integer value specifying count for the number of keys generated for the session_id.

H2OCoxPHModel-class

The H2OCoxPHModel object.

Description
Virtual object representing H2O's CoxPH Model.

Usage

## S4 method for signature 'H2OCoxPHModel'
show(object)

## S3 method for class 'H2OCoxPHModel'
coef(object, ...)

## S3 method for class 'H2OCoxPHModel'
extractAIC(fit, scale, k = 2, ...)

## S3 method for class 'H2OCoxPHModel'
logLik(object, ...)

## S3 method for class 'H2OCoxPHModel'
survfit.H2OCoxPHModel(formula, newdata, ...)

## S3 method for class 'H2OCoxPHModel'
vcov(object, ...)

Arguments

object  an H2OCoxPHModel object.

...  additional arguments to pass on.

fit  an H2OCoxPHModel object.
scale
k
formula
newdata

optional numeric specifying the scale parameter of the model.
numeric specifying the weight of the equivalent degrees of freedom.
an H2OCoxPHModel object.
an optional H2OFrame or data.frame with the same variable names as those that appear in the H2OCoxPHModel object.

H2OCoxPHModelSummary-class

The H2OCoxPHModelSummary object.

Description
Wrapper object for summary information compatible with survival package.

Usage

## S4 method for signature 'H2OCoxPHModelSummary'
show(object)

## S3 method for class 'H2OCoxPHModelSummary'
coef(object, ...)

Arguments

object An H2OCoxPHModelSummary object.
... additional arguments to pass on.

Slots

summary A list containing the a summary compatible with CoxPH summary used in the survival package.

H2OFrame-class

The H2OFrame class

Description

This class represents an H2OFrame object
Description

Operators to extract or replace parts of H2OFrame objects.

Usage

```r
# S3 method for class 'H2OFrame'
data[row, col, drop = TRUE]

# S3 method for class 'H2OFrame'
x$name

# S3 method for class 'H2OFrame'
x[[i, exact = TRUE]]

# S3 method for class 'H2OFrame'
x$name

# S3 method for class 'H2OFrame'
x[[i, exact = TRUE]]

# S3 replacement method for class 'H2OFrame'
data[row, col, ...] <- value

# S3 replacement method for class 'H2OFrame'
data$name <- value

# S3 replacement method for class 'H2OFrame'
data[[name]] <- value
```

Arguments

- `data` object from which to extract element(s) or in which to replace element(s).
- `row` index specifying row element(s) to extract or replace. Indices are numeric or character vectors or empty (missing) or will be matched to the names.
- `col` index specifying column element(s) to extract or replace.
- `drop` Unused
- `x` An H2OFrame
- `name` a literal character string or a name (possibly backtick quoted).
- `i` index
- `exact` controls possible partial matching of `[[` when extracting a character
- `...` Further arguments passed to or from other methods.
H2OGrid-class

value To be assigned

---

**H2OGrid-class**

**H2O Grid**

**Description**

A class to contain the information about grid results

**Usage**

```r
## S4 method for signature 'H2OGrid'
show(object)
```

**Arguments**

- `object` an H2OGrid object.

**Slots**

- `grid_id` the final identifier of grid
- `model_ids` list of model IDs which are included in the grid object
- `hyper_names` list of parameter names used for grid search
- `failed_params` list of model parameters which caused a failure during model building, it can contain a null value
- `failure_details` list of detailed messages which correspond to failed parameters field
- `failure_stack_traces` list of stack traces corresponding to model failures reported by failed_params and failure_details fields
- `failed_raw_params` list of failed raw parameters
- `summary_table` table of models built with parameters and metric information.

**See Also**

[H2OModel](#) for the final model types.
## H2OInfogram-class

### Description

wrapper function for instantiating H2OInfogram

### Usage

```r
H2OInfogram(model_id, ...)  
```

### Arguments

- `model_id`: is string of H2OModel object
- `...`: parameters to algorithm, admissible_features, ...

### Value

A H2OInfogram object

## H2OInfogram-class

### Description

H2OInfogram class contains a subset of what a normal H2OModel will return

### Slots

- `model_id`: string returned as part of every H2OModel
- `algorithm`: string denoting the algorithm used to build infogram
- `admissible_features`: string array denoting all predictor names which pass the cmi and relevance threshold
- `admissible_features_valid`: string array denoting all predictor names which pass the cmi and relevance threshold from validation frame
- `admissible_features_xval`: string array denoting all predictor names which pass the cmi and relevance threshold from cv holdout set
- `net_information_threshold`: numeric value denoting threshold used for predictor selection
- `total_information_threshold`: numeric value denoting threshold used for predictor selection
- `safety_index_threshold`: numeric value denoting threshold used for predictor selection
- `relevance_index_threshold`: numeric value denoting threshold used for predictor selection
- `admissible_score`: H2OFrame that contains columns, admissible, admissible_index, relevance, cmi, cmi_raw
admissible_score_valid H2OFrame that contains columns, admissible, admissible_index, relevance, cmi, cmi_raw from validation frame

admissible_score_xval H2OFrame that contains averages of columns, admissible, admissible_index, relevance, cmi, cmi_raw from cv hold-out

H2OLeafNode-class  The H2OLeafNode class.

Description

This class represents a single leaf node in an H2OTree.

Details

#' @aliases H2OLeafNode

H2OModel-class  The H2OModel object.

Description

This virtual class represents a model built by H2O.

Usage

## S4 method for signature 'H2OModel'
show(object)

Arguments

object  an H2OModel object.

Details

This object has slots for the key, which is a character string that points to the model key existing in the H2O cluster, the data used to build the model (an object of class H2OFrame).
H2OModelMetrics-class

Slots

model_id A character string specifying the key for the model fit in the H2O cluster’s key-value store.

algorithm A character string specifying the algorithm that were used to fit the model.

parameters A list containing the parameter settings that were used to fit the model that differ from the defaults.

allparameters A list containing all parameters used to fit the model.

have_pojo A logical indicating whether export to POJO is supported

have_mojo A logical indicating whether export to MOJO is supported

model A list containing the characteristics of the model returned by the algorithm.

H2OModelFuture-class H2O Future Model

Description

A class to contain the information for background model jobs.

Slots

job_key a character key representing the identification of the job process.

model_id the final identifier for the model

See Also

H2OModel for the final model types.

H2OModelMetrics-class The H2OModelMetrics Object.

Description

A class for constructing performance measures of H2O models.
Usage

```r
## S4 method for signature 'H2OModelMetrics'
show(object)

## S4 method for signature 'H2OBinomialMetrics'
show(object)

## S4 method for signature 'H2OBinomialUpliftMetrics'
show(object)

## S4 method for signature 'H2OMultinomialMetrics'
show(object)

## S4 method for signature 'H2OOrdinalMetrics'
show(object)

## S4 method for signature 'H2ORegressionMetrics'
show(object)

## S4 method for signature 'H2OClusteringMetrics'
show(object)

## S4 method for signature 'H2OAutoEncoderMetrics'
show(object)

## S4 method for signature 'H2ODimReductionMetrics'
show(object)

## S4 method for signature 'H2OAnomalyDetectionMetrics'
show(object)
```

Arguments

- `object` An `H2OModelMetrics` object

Description

The `H2ONode-class` represents the H2O Node class.

Usage

```r
## S4 method for signature 'H2ONode'
show(object)
```
Arguments

object  an H2ONode object.

Slots

id  An integer representing node's unique identifier. Generated by H2O.
levels  A character representing categorical levels on split from parent's node belonging into this node. NULL for root node or non-categorical splits.

# @aliases H2ONode

H2OSegmentModels-class

H2O Segment Models

Description

A class to contain the information for segment models.

Usage

## S4 method for signature 'H2OSegmentModels'
show(object)

Arguments

object  an H2OModel object.

Slots

segment_models_id  the identifier for the segment models collections

H2OSegmentModelsFuture-class

H2O Future Segment Models

Description

A class to contain the information for background segment models jobs.

Slots

job_key  a character key representing the identification of the job process.
segment_models_id  the final identifier for the segment models collections

See Also

H2OSegmentModels for the final segment models types.
**H2OSplitNode-class**

*The H2OSplitNode class.*

**Description**

This class represents a single non-terminal node in an H2OTree.

**Slots**

- **threshold** A numeric split threshold, typically when the split column is numerical.
- **left_child** A H2ONodeOrNULL representing the left child node, if a node has one.
- **right_child** A H2ONodeOrNULL representing the right child node, if a node has one.
- **split_feature** A character representing the name of the column this node splits on.
- **left_levels** A character representing the levels of a categorical feature heading to the left child of this node. NA for non-categorical split.
- **right_levels** A character representing the levels of a categorical feature heading to the right child of this node. NA for non-categorical split.
- **na_direction** A character representing the direction of NA values. **LEFT** means NA values go to the left child node, **RIGH** means NA values go to the right child node.

**H2OTree-class**

*The H2OTree class.*

**Description**

This class represents a model of a Tree built by one of H2O’s algorithms (GBM, Random Forest).

**Usage**

```r
## S4 method for signature 'H2OTree'
show(object)
```

**Arguments**

- **object** an H2OTree object.

**Slots**

- **root_node** A H2ONode representing the beginning of the tree behind the model. Allows further tree traversal.
- **left_children** An integer vector with left child nodes of tree’s nodes
- **right_children** An integer vector with right child nodes of tree’s nodes
node_ids  An integer representing identification number of a node. Node IDs are generated by H2O.

descriptions A character vector with descriptions for each node to be found in the tree. Contains split threshold if the split is based on numerical column. For categorical splits, it contains list of categorical levels for transition from the parent node.

model_id  A character with the name of the model this tree is related to.

tree_number An integer representing the order in which the tree has been built in the model.

tree_class A character representing name of tree’s class. Number of tree classes equals to the number of levels in categorical response column. As there is exactly one class per categorical level, name of tree’s class equals to the corresponding categorical level of response column. In case of regression and binomial, the name of the categorical level is ignored can be omitted, as there is exactly one tree built in both cases.

thresholds  A numeric split thresholds. Split thresholds are not only related to numerical splits, but might be present in case of categorical split as well.

features A character with names of the feature/column used for the split.

levels A character representing categorical levels on split from parent’s node belonging into this node. NULL for root node or non-categorical splits.

nas  A character representing if NA values go to the left node or right node. May be NA if node is a leaf.

predictions  A numeric representing predictions for each node in the graph.

tree_decision_path A character, plain language rules representation of a trained decision tree

decision_paths A character representing plain language rules that were used in a particular prediction.

left_cat_split A character list of categorical levels leading to the left child node. Only present when split is categorical, otherwise none.

right_cat_split A character list of categorical levels leading to the right child node. Only present when split is categorical, otherwise none.

---

**housevotes**

*United States Congressional Voting Records 1984*

**Description**

This data set includes votes for each of the U.S. House of Representatives Congressmen on the 16 key votes identified by the CQA. The CQA lists nine different types of votes: voted for, paired for, and announced for (these three simplified to yea), voted against, paired against, and announced against (these three simplified to nay), voted present, voted present to avoid conflict of interest, and did not vote or otherwise make a position known (these three simplified to an unknown disposition).

**Format**

A data frame with 435 rows and 17 columns
Source

References

Description
Method on H2OInfogram object which in this case is to instantiate and initialize it

Usage
```r
## S4 method for signature 'H2OInfogram'
initialize(.Object, model_id, ...)
```

Arguments

- `.Object` An H2OInfogram object
- `model_id` string returned as part of every H2OModel
- `...` additional arguments to pass on

Value
A H2OInfogram object

iris

Edgar Anderson’s Iris Data

Description
Measurements in centimeters of the sepal length and width and petal length and width, respectively, for three species of iris flowers.

Format
A data frame with 150 rows and 5 columns
Source


The data were collected by Anderson, Edgar (1935). The irises of the Gaspe Peninsula, Bulletin of the American Iris Society, 59, 2-5.

is.character

Check if character

Description

Check if character

Usage

is.character(x)

Arguments

x

An H2OFrame object

Examples

## Not run:
library(h2o)
h2o.init()

f <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/coxph_test/heart.csv"
heart <- h2o.importFile(f)

heart["transplant"] <- as.character(heart["transplant"])
is.character(heart["transplant"])

## End(Not run)

is.factor

Check if factor

Description

Check if factor

Usage

is.factor(x)
is.h2o

Arguments

x  An H2OFrame object

Description
Test if object is H2O Frame.

Usage

is.h2o(x)

Arguments

x  An R object.

Examples

## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)

is.h2o(frame)

## End(Not run)

is.numeric

Description
Check if numeric

Usage

is.numeric(x)

Arguments

x  An H2OFrame object
### Keyed-class

**Virtual Keyed class**

**Description**

Base class for all objects having a persistent representation on backend.

**length,H2OTree-method**

*Overrides the behavior of length() function on H2OTree class. Returns number of nodes in an H2OTree*

**Description**

Overrides the behavior of length() function on H2OTree class. Returns number of nodes in an H2OTree

**Usage**

```r
## S4 method for signature 'H2OTree'
length(x)
```

**Arguments**

- `x` An H2OTree to count nodes for.

### Logical-or

**Logical or for H2OFrames**

**Description**

Logical or for H2OFrames

**Usage**

```
`||`(x, y)
```

**Arguments**

- `x` An H2OFrame object
- `y` An H2OFrame object
ModelAccessors

Accessor Methods for H2OModel Object

Description

Function accessor methods for various H2O output fields.

Usage

getParms(object)

## S4 method for signature 'H2OModel'
getParms(object)

getCenters(object)

getCentersStd(object)

getWithinSS(object)

getTotWithinSS(object)

getBetweenSS(object)

getTotSS(object)

getIterations(object)

getClusterSizes(object)

## S4 method for signature 'H2OClusteringModel'
getCenters(object)

## S4 method for signature 'H2OClusteringModel'
getCentersStd(object)

## S4 method for signature 'H2OClusteringModel'
getWithinSS(object)

## S4 method for signature 'H2OClusteringModel'
getTotWithinSS(object)

## S4 method for signature 'H2OClusteringModel'
getBetweenSS(object)

## S4 method for signature 'H2OClusteringModel'
getTotSS(object)
## S4 method for signature 'H2OClusteringModel'
getIterations(object)

## S4 method for signature 'H2OClusteringModel'
getClusterSizes(object)

### Arguments

- **object**: an **H2OModel** class object.

---

model_cache-class

---

*Needed to be able to memoise the models*

---

### Description

Needed to be able to memoise the models

---

### names.H2OFrame

*Column names of an H2OFrame*

---

### Description

Column names of an H2OFrame

### Usage

```r
## S3 method for class 'H2OFrame'
names(x)
```

### Arguments

- **x**: An H2OFrame

### Examples

```r
## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)

names(frame)

## End(Not run)
```
**Description**

Methods for group generic functions and H2O objects.

**Usage**

```r
## S3 method for class 'H2OFrame'
Ops(e1, e2)

## S3 method for class 'H2OFrame'
Math(x, ...)  

## S3 method for class 'H2OFrame'
Summary(x, ..., na.rm)

## S3 method for class 'H2OFrame'
!x

## S3 method for class 'H2OFrame'
is.na(x)

## S3 method for class 'H2OFrame'
t(x)

log(x, ...)

log10(x)

log2(x)

log1p(x)

trunc(x, ...)

x %% y

nrow.H2OFrame(x)
```
ncol.H2OFrame(x)

## S3 method for class 'H2OFrame'
length(x)

h2o.length(x)

## S3 replacement method for class 'H2OFrame'
names(x) <- value

colnames(x) <- value

Arguments

e1 object
e2 object
x object
... Further arguments passed to or from other methods.
na.rm logical. whether or not missing values should be removed
y object
value To be assigned

---

plot.H2OInfogram  Plot an H2O Infogram

Description

Plots the Infogram for an H2OInfogram object.

Usage

## S3 method for class 'H2OInfogram'
plot(x, ...)

Arguments

x A fitted H2OInfogram object.
... additional arguments to pass on.

Value

A ggplot2 object.

See Also

h2o.infogram
Examples

```r
## Not run:
  h2o.init()

  # Convert iris dataset to an H2OFrame
  train <- as.h2o(iris)

  # Create and plot infogram
  ig <- h2o.infogram(y = "Species", training_frame = train)
  plot(ig)

  ## End(Not run)
```

plot.H2OModel

Plot an H2O Model

Description

Plots training set (and validation set if available) scoring history for an H2O Model

Usage

```r
## S3 method for class 'H2OModel'
plot(x, timestep = "AUTO", metric = "AUTO", ...)
```

Arguments

- `x`: A fitted `H2OModel` object for which the scoring history plot is desired.
- `timestep`: A unit of measurement for the x-axis.
- `metric`: A unit of measurement for the y-axis.
- `...`: additional arguments to pass on.

Details

This method dispatches on the type of H2O model to select the correct scoring history. The timestep and metric arguments are restricted to what is available in the scoring history for a particular type of model.

Value

Returns a scoring history plot.

See Also

- `h2o.deeplearning`, `h2o.gbm`, `h2o.glm`, `h2o.randomForest` for model generation in h2o.
Examples

### Not run:

```r
if (requireNamespace("mlbench", quietly=TRUE)) {
  library(h2o)
  h2o.init()

  df <- as.h2o(mlbench::mlbench.friedman1(10000, 1))
  rng <- h2o.runif(df, seed = 1234)
  train <- df[rng < 0.8,]
  valid <- df[rng >= 0.8,]

  gbm <- h2o.gbm(x = 1:10, y = "y", training_frame = train, validation_frame = valid,
                  ntrees = 500, learn_rate = 0.01, score_each_iteration = TRUE)
  plot(gbm)
  plot(gbm, timestep = "duration", metric = "deviance")
  plot(gbm, timestep = "number_of_trees", metric = "deviance")
  plot(gbm, timestep = "number_of_trees", metric = "rmse")
  plot(gbm, timestep = "number_of_trees", metric = "mae")
}
### End(Not run)
```

---

**plot.H2OTabulate**

*Plot an H2O Tabulate Heatmap*

**Description**

Plots the simple co-occurrence based tabulation of X vs Y as a heatmap, where X and Y are two Vecs in a given dataset. This function requires suggested ggplot2 package.

**Usage**

```r
## S3 method for class 'H2OTabulate'
plot(x, xlab = x$cols[1], ylab = x$cols[2], base_size = 12, ...)
```

**Arguments**

- `x`  
  An H2OTabulate object for which the heatmap plot is desired.
- `xlab`  
  A title for the x-axis. Defaults to what is specified in the given H2OTabulate object.
- `ylab`  
  A title for the y-axis. Defaults to what is specified in the given H2OTabulate object.
- `base_size`  
  Base font size for plot.
- `...`  
  Additional arguments to pass on.

**Value**

Returns a ggplot2-based heatmap of co-occurrence.
predict.H2OAutoML

See Also

h2o.tabulate

Examples

```r
## Not run:
library(h2o)
h2o.init()
df <- as.h2o(iris)
tab <- h2o.tabulate(data = df, x = "Sepal.Length", y = "Petal.Width",
                   weights_column = NULL, nbins_x = 10, nbins_y = 10)
plot(tab)

## End(Not run)
```

predict.H2OAutoML  Predict on an AutoML object

Description

Obtains predictions from an AutoML object.

Usage

```r
## S3 method for class 'H2OAutoML'
predict(object, newdata, ...)

## S3 method for class 'H2OAutoML'
h2o.predict(object, newdata, ...)
```

Arguments

- `object` a fitted `H2OAutoML` object for which prediction is desired
- `newdata` An H2OFrame object in which to look for variables with which to predict.
- `...` additional arguments to pass on.

Details

This method generated predictions on the leader model from an AutoML run. The order of the rows in the results is the same as the order in which the data was loaded, even if some rows fail (for example, due to missing values or unseen factor levels).

Value

Returns an H2OFrame object with probabilities and default predictions.
predict.H2OModel

Predict on an H2O Model

Description

Obtains predictions from various fitted H2O model objects.

Usage

## S3 method for class 'H2OModel'
predict(object, newdata, ...)

## S3 method for class 'H2OModel'
h2o.predict(object, newdata, ...)

Arguments

- **object**: a fitted H2OModel object for which prediction is desired
- **newdata**: An H2OFrame object in which to look for variables with which to predict.
- **...**: additional arguments to pass on.

Details

This method dispatches on the type of H2O model to select the correct prediction/scoring algorithm. The order of the rows in the results is the same as the order in which the data was loaded, even if some rows fail (for example, due to missing values or unseen factor levels).

Value

Returns an H2OFrame object with probabilities and default predictions.

See Also

h2o.deeplearning, h2o.gbm, h2o.glm, h2o.randomForest for model generation in h2o.

Examples

```r
## Not run:
library(h2o)
h2o.init()

f <- "https://s3.amazonaws.com/h2o-public-test-data/smalldata/glm_test/insurance.csv"
insurance <- h2o.importFile(f)
predictors <- colnames(insurance)[1:4]
response <- "Claims"
insurance["Group"] <- as.factor(insurance["Group"])
insurance["Age"] <- as.factor(insurance["Age"])
splits <- h2o.splitFrame(data = insurance, ratios = 0.8, seed = 1234)
```
train <- splits[[1]]
valid <- splits[[2]]
insurance_gbm <- h2o.gbm(x = predictors, y = response,
                        training_frame = train,
                        validation_frame = valid,
                        distribution = "huber",
                        huber_alpha = 0.9, seed = 1234)
h2o.predict(insurance_gbm, newdata = insurance)

## End(Not run)

**predict_contributions.H2OModel**

Predict feature contributions - SHAP values on an H2O Model (only DRF, GBM, XGBoost models and equivalent imported MOJOs).

**Description**

Default implementation return H2OFrame shape (#rows, #features + 1) - there is a feature contribution column for each input feature, the last column is the model bias (same value for each row). The sum of the feature contributions and the bias term is equal to the raw prediction of the model. Raw prediction of tree-based model is the sum of the predictions of the individual trees before the inverse link function is applied to get the actual prediction. For Gaussian distribution the sum of the contributions is equal to the model prediction.

**Usage**

```r
predict_contributions.H2OModel(
    object,
    newdata,
    output_format = c("original", "compact"),
    top_n = 0,
    bottom_n = 0,
    compare_abs = FALSE,
    ...
)
```

```r
h2o.predict_contributions(
    object,
    newdata,
    output_format = c("original", "compact"),
    top_n = 0,
    bottom_n = 0,
    compare_abs = FALSE,
    ...
)
```
predict_contributions.H2OModel

Arguments

- **object**: a fitted H2OModel object for which prediction is desired
- **newdata**: An H2OFrame object in which to look for variables with which to predict.
- **output_format**: Specify how to output feature contributions in XGBoost - XGBoost by default outputs contributions for 1-hot encoded features, specifying a compact output format will produce a per-feature contribution. Defaults to original.
- **top_n**: Return only #top_n highest contributions + bias If top_n<0 then sort all SHAP values in descending order If top_n<0 then sort all SHAP values in descending order
- **bottom_n**: Return only #bottom_n lowest contributions + bias If top_n and bottom_n are defined together then return array of #top_n + #bottom_n + bias If bottom_n<0 then sort all SHAP values in ascending order If top_n<0 then sort all SHAP values in descending order
- **compare_abs**: True to compare absolute values of contributions
- ... additional arguments to pass on.

Details

Note: Multinomial classification models are currently not supported.

Value

Returns an H2OFrame contain feature contributions for each input row.

See Also

h2o.gbm and h2o.randomForest for model generation in h2o.

Examples

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
prostate_gbm <- h2o.gbm(3:9, "AGE", prostate)
h2o.predict(prostate_gbm, prostate)
# Compute SHAP
h2o.predict_contributions(prostate_gbm, prostate)
# Compute SHAP and pick the top two highest
h2o.predict_contributions(prostate_gbm, prostate, top_n=2)
# Compute SHAP and pick the top two lowest
h2o.predict_contributions(prostate_gbm, prostate, bottom_n=2)
# Compute SHAP and pick the top two highest regardless of the sign
h2o.predict_contributions(prostate_gbm, prostate, top_n=2, compare_abs=TRUE)
# Compute SHAP and pick the top two lowest regardless of the sign
h2o.predict_contributions(prostate_gbm, prostate, bottom_n=2, compare_abs=TRUE)
# Compute SHAP values and show them all in descending order
h2o.predict_contributions(prostate_gbm, prostate, top_n=-1)
```
predict_leaf_node_assignment.H2OModel(427)

# Compute SHAP and pick the top two highest and top two lowest
h2o.predict_contributions(prostate_gbm, prostate, top_n=2, bottom_n=2)

## End(Not run)

---

**predict_leaf_node_assignment.H2OModel**

*Predict the Leaf Node Assignment on an H2O Model*

---

**Description**

Obtains leaf node assignment from fitted H2O model objects.

**Usage**

```r
predict_leaf_node_assignment.H2OModel(
  object,
  newdata,
  type = c("Path", "Node_ID"),
  ...
)
```

```r
h2o.predict_leaf_node_assignment(
  object,
  newdata,
  type = c("Path", "Node_ID"),
  ...
)
```

**Arguments**

- **object**: a fitted `H2OModel` object for which prediction is desired
- **newdata**: An H2OFrame object in which to look for variables with which to predict.
- **type**: choice of either "Path" when tree paths are to be returned (default); or "Node_ID" when the output
- **...**: additional arguments to pass on.

**Details**

For every row in the test set, return the leaf placements of the row in all the trees in the model. Placements can be represented either by paths to the leaf nodes from the tree root or by H2O’s internal identifiers. The order of the rows in the results is the same as the order in which the data was loaded.

**Value**

Returns an H2OFrame object with categorical leaf assignment identifiers for each tree in the model.
See Also

h2o.gbm and h2o.randomForest for model generation in h2o.

Examples

## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
prostate$CAPSULE <- as.factor(prostate$CAPSULE)
prostate_gbm <- h2o.gbm(3:9, "CAPSULE", prostate)
h2o.predict(prostate_gbm, prostate)
h2o.predict_leaf_node_assignment(prostate_gbm, prostate)

## End(Not run)
print.H2OTable

## End(Not run)

### print.H2OTable

*Print method for H2OTable objects*

**Description**

This will print a truncated view of the table if there are more than 20 rows.

**Usage**

```r
## S3 method for class 'H2OTable'
print(x, header = TRUE, ...)
```

**Arguments**

- `x`: An H2OTable object
- `header`: A logical value dictating whether or not the table name should be printed.
- `...`: Further arguments passed to or from other methods.

**Value**

The original `x` object

**Examples**

```r
## Not run:
library(h2o)
h2o.init()

cars <- h2o.importFile(f)
print(cars, header = TRUE)

## End(Not run)
```
prostate  Prostate Cancer Study

Description
Baseline exam results on prostate cancer patients from Dr. Donn Young at The Ohio State University Comprehensive Cancer Center.

Format
A data frame with 380 rows and 9 columns

Source

range.H2OFrame  Range of an H2O Column

Description
Range of an H2O Column

Usage
## S3 method for class 'H2OFrame'
range(..., na.rm = TRUE)

Arguments
...  An H2OFrame object.
na.rm  ignore missing values

Examples
## Not run:
library(h2o)
h2o.init()

frame <- h2o.createFrame(rows = 6, cols = 2,
categorical_fraction = 0.0,
missing_fraction = 0.7,
seed = 123)

range(frame, na.rm = TRUE)

## End(Not run)
Scaling and Centering of an H2OFrame

Description
Centers and/or scales the columns of an H2O dataset.

Usage

```r
## S3 method for class 'H2OFrame'
scale(x, center = TRUE, scale = TRUE)
```

Arguments
- `x`: An H2OFrame object.
- `center`: either a logical value or numeric vector of length equal to the number of columns of `x`.
- `scale`: either a logical value or numeric vector of length equal to the number of columns of `x`.

Examples

```r
## Not run:
library(h2o)
h2o.init()
iris_hf <- as.h2o(iris)
summary(iris_hf)

# Scale and center all the numeric columns in iris data set
iris_scaled <- scale(iris_hf[, 1:4])
## End(Not run)
```

Format AutoML object in user-friendly way

Description
Format AutoML object in user-friendly way

Usage

```r
## S4 method for signature 'H2OAutoML'
show(object)
```

Arguments
- `object`: an H2OAutoML object.
staged_predict_proba.H2OModel

Predict class probabilities at each stage of an H2O Model

Description

The output structure is analogous to the output of `h2o.predict_leaf_node_assignment`. For each tree \( t \) and class \( c \) there will be a column \( T_t.C_c \) (e.g., \( T_3.C_1 \) for tree 3 and class 1). The value will be the corresponding predicted probability of this class by combining the raw contributions of trees \( T_1.C_c,...,T_t.C_c \). Binomial models build the trees just for the first class and values in columns \( T_x.C_1 \) thus correspond to the probability \( p_0 \).

Usage

```r
staged_predict_proba.H2OModel(object, newdata, ...)

h2o.staged_predict_proba(object, newdata, ...)
```

Arguments

- `object` a fitted `H2OModel` object for which prediction is desired
- `newdata` An `H2OFrame` object in which to look for variables with which to predict.
- `...` additional arguments to pass on.

Value

Returns an `H2OFrame` object with predicted probability for each tree in the model.

See Also

`h2o.gbm` and `h2o.randomForest` for model generation in `h2o`.

Examples

```r
## Not run:
library(h2o)
h2o.init()
prostate_path <- system.file("extdata", "prostate.csv", package = "h2o")
prostate <- h2o.uploadFile(path = prostate_path)
prostate$CAPSULE <- as.factor(prostate$CAPSULE)
prostate_gbm <- h2o.gbm(3:9, "CAPSULE", prostate)
h2o.predict(prostate_gbm, prostate)
h2o.staged_predict_proba(prostate_gbm, prostate)

## End(Not run)
```
### str.H2OFrame

**Description**

Display the structure of an H2OFrame object

**Usage**

```r
## S3 method for class 'H2OFrame'
str(object, ..., cols = FALSE)
```

**Arguments**

- `object` An H2OFrame.
- `...` Further arguments to be passed from or to other methods.
- `cols` Print the per-column str for the H2OFrame

### summary,H2OAutoML-method

**Description**

Format AutoML object in user-friendly way

**Usage**

```r
## S4 method for signature 'H2OAutoML'
summary(object)
```

**Arguments**

- `object` an H2OAutoML object.
Summary method for H2OCoxPHModel objects

**Description**

Summary method for H2OCoxPHModel objects

**Usage**

```r
## S4 method for signature 'H2OCoxPHModel'
summary(object, conf.int = 0.95, scale = 1)
```

**Arguments**

- `object`: an H2OCoxPHModel object.
- `conf.int`: a specification of the confidence interval.
- `scale`: a scale.

Format grid object in user-friendly way

**Description**

Format grid object in user-friendly way

**Usage**

```r
## S4 method for signature 'H2OGrid'
summary(object, show_stack_traces = FALSE)
```

**Arguments**

- `object`: an H2OGrid object.
- `show_stack_traces`: a flag to show stack traces for model failures
summary.H2OModel-method

Print the Model Summary

Description

Print the Model Summary

Usage

## S4 method for signature 'H2OModel'
summary(object, ...)

Arguments

object An H2OModel object.
...

further arguments to be passed on (currently unimplemented)

use.package

Use optional package

Description

Testing availability of optional package, its version, and extra global default. This function is used internally. It is exported and documented because user can control behavior of the function by global option.

Usage

use.package(
  package,
  version = "1.9.8"[package == "data.table"],
  use = getOption("h2o.use.data.table", TRUE)[package == "data.table"]
)

Arguments

package character scalar name of a package that we Suggests or Enhances on.
version character scalar required version of a package.
use logical scalar, extra escape option, to be used as global option.
Details

We use this function to control csv read/write with optional data.table package. Currently data.table is enabled by default for some operations, to disable it set options("h2o.use.data.table"=FALSE). It is possible to control just fread or fwrite with options("h2o.fread"=FALSE,"h2o.fwrite"=FALSE). h2o.fread and h2o.fwrite options are not handled in this function but next to fread and fwrite calls.

See Also

as.h2o.data.frame, as.data.frame.H2OFrame

Examples

op <- options("h2o.use.data.table" = TRUE)
if (use.package("data.table")) {
  cat("optional package data.table 1.9.8+ is available\n")
} else {
  cat("optional package data.table 1.9.8+ is not available\n")
}
options(op)

---

walking Muscular Actuations for Walking Subject

Description

The musculoskeletal model, experimental data, settings files, and results for three-dimensional, muscle-actuated simulations at walking speed as described in Hamner and Delp (2013). Simulations were generated using OpenSim 2.4. The data is available from https://simtk.org/frs/index.php?group_id=603.

Format

A data frame with 151 rows and 124 columns

References

with_no_h2o_progress  Suppresses h2o progress output from expr

Description

Suppresses h2o progress output from expr

Usage

with_no_h2o_progress(expr)

Arguments

expr  expression

Value

result of expr

zzz  Shutdown H2O cluster after examples run

Description

Shutdown H2O cluster after examples run

Examples

## Not run:
library(h2o)
h2o.init()
h2o.shutdown(prompt = FALSE)
Sys.sleep(3)

## End(Not run)
Logical and for H2OFrames

Description

Logical and for H2OFrames

Usage

'& & '(x, y)

Arguments

x    An H2OFrame object
y    An H2OFrame object
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