Package 'hardhat'

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Title Construct Modeling Packages

Version 1.0.0

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Description Building modeling packages is hard. A large amount of effort generally goes into providing an implementation for a new method that is efficient, fast, and correct, but often less emphasis is put on the user interface. A good interface requires specialized knowledge about S3 methods and formulas, which the average package developer might not have. The goal of 'hardhat' is to reduce the burden around building new modeling packages by providing functionality for preprocessing, predicting, and validating input.

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URL https://github.com/tidymodels/hardhat, https://hardhat.tidymodels.org

 $\mathbf{BugReports}\ \mathsf{https://github.com/tidymodels/hardhat/issues}$

```
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tibble (i=3.1.7),
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```

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 ${\bf Config/Needs/website}\ {\it tidyverse/tidytemplate}$

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LazyData true

Roxygen list(markdown = TRUE)

61

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${\bf Config/test that/edition} \ \ 3$

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modeling-package

Create a modeling package

Description

create_modeling_package() will:

- Call usethis::create_package() to set up a new R package.
- Call use_modeling_deps().
- Call use_modeling_files().

use_modeling_deps() will:

- Add hardhat, rlang, and stats to Imports
- Add recipes to Suggests
- If roxygen2 is available, use roxygen markdown

use_modeling_files() will:

- Add a package documentation file
- Generate and populate 3 files in R/:

```
-~\{\{\mathsf{model}\}\}\text{-}\mathsf{constructor.R}
```

- {{model}}-fit.R
- {{model}}-predict.R

Usage

```
create_modeling_package(path, model, fields = NULL, open = interactive())
use_modeling_deps()
use_modeling_files(model)
```

Arguments

path

A path. If it exists, it is used. If it does not exist, it is created, provided that the parent path exists.

model

A string. The name of the high level modeling function that users will call. For example, "linear_regression". This will be used to populate the skeleton. Spaces are not allowed.

fields

A named list of fields to add to DESCRIPTION, potentially overriding default values. See usethis::use_description() for how you can set personalized defaults using package options.

open

If TRUE, activates the new project:

- If RStudio desktop, the package is opened in a new session.
- $\bullet\,$ If on RS tudio server, the current RS tudio project is activated.
- Otherwise, the working directory and active project is changed.

Value

```
create_modeling_package() returns the project path invisibly.
use_modeling_deps() returns invisibly.
use_modeling_files() return model invisibly.
```

add_intercept_column

Add an intercept column to data

Description

This function adds an integer column of 1's to data.

Usage

```
add_intercept_column(data, name = "(Intercept)")
```

Arguments

data A data frame or matrix.

name The name for the intercept column. Defaults to "(Intercept)", which is

the same name that stats::lm() uses.

Details

If a column named name already exists in data, then data is returned unchanged and a warning is issued.

Value

data with an intercept column.

```
add_intercept_column(mtcars)
add_intercept_column(mtcars, "intercept")
add_intercept_column(as.matrix(mtcars))
```

```
default_formula_blueprint
```

Default formula blueprint

Description

This pages holds the details for the formula preprocessing blueprint. This is the blueprint used by default from mold() if x is a formula.

Usage

```
default_formula_blueprint(
  intercept = FALSE,
  allow_novel_levels = FALSE,
  indicators = "traditional",
  composition = "tibble"
)

## S3 method for class 'formula'
mold(formula, data, ..., blueprint = NULL)
```

Arguments

intercept

A logical. Should an intercept be included in the processed data? This information is used by the process function in the mold and forge function list.

allow_novel_levels

A logical. Should novel factor levels be allowed at prediction time? This information is used by the clean function in the forge function list, and is passed on to scream().

indicators

A single character string. Control how factors are expanded into dummy variable indicator columns. One of:

- "traditional" The default. Create dummy variables using the traditional model.matrix() infrastructure. Generally this creates K
 -1 indicator columns for each factor, where K is the number of levels in that factor.
- "none" Leave factor variables alone. No expansion is done.
- "one_hot" Create dummy variables using a one-hot encoding approach that expands unordered factors into all K indicator columns, rather than K -1.

 ${\tt composition}$

Either "tibble", "matrix", or "dgCMatrix" for the format of the processed predictors. If "matrix" or "dgCMatrix" are chosen, all of the predictors must be numeric after the preprocessing method has been applied; otherwise an error is thrown.

formula

A formula specifying the predictors and the outcomes.

data

A data frame or matrix containing the outcomes and predictors.

. . .

Not used.

blueprint

A preprocessing blueprint. If left as NULL, then a $default_formula_blueprint()$ is used.

Details

While not different from base R, the behavior of expanding factors into dummy variables when indicators = "traditional" and an intercept is *not* present is not always intuitive and should be documented.

- When an intercept is present, factors are expanded into K-1 new columns, where K is the number of levels in the factor.
- When an intercept is *not* present, the first factor is expanded into all K columns (one-hot encoding), and the remaining factors are expanded into K-1 columns. This behavior ensures that meaningful predictions can be made for the reference level of the first factor, but is not the exact "no intercept" model that was requested. Without this behavior, predictions for the reference level of the first factor would always be forced to 0 when there is no intercept.

Offsets can be included in the formula method through the use of the inline function stats::offset(). These are returned as a tibble with 1 column named ".offset" in the \$extras\$offset slot of the return value.

Value

For default_formula_blueprint(), a formula blueprint.

Mold

When mold() is used with the default formula blueprint:

• Predictors

- The RHS of the formula is isolated, and converted to its own 1 sided formula: $\tilde{\ }$ RHS.
- Runs stats::model.frame() on the RHS formula and uses data.
- If indicators = "traditional", it then runs stats::model.matrix() on the result.
- If indicators = "none", factors are removed before model.matrix() is run, and then added back afterwards. No interactions or inline functions involving factors are allowed.
- If indicators = "one_hot", it then runs stats::model.matrix() on the result using a contrast function that creates indicator columns for all levels of all factors.
- If any offsets are present from using offset(), then they are extracted with model_offset().
- If intercept = TRUE, adds an intercept column.
- Coerces the result of the above steps to a tibble.

Outcomes

- The LHS of the formula is isolated, and converted to its own 1 sided formula: ~
- Runs stats::model.frame() on the LHS formula and uses data.
- Coerces the result of the above steps to a tibble.

Forge

When forge() is used with the default formula blueprint:

- It calls shrink() to trim new_data to only the required columns and coerce new_data to a tibble.
- It calls scream() to perform validation on the structure of the columns of new_data.
- Predictors
 - It runs stats::model.frame() on new_data using the stored terms object corresponding to the predictors.
 - If, in the original mold() call, indicators = "traditional" was set, it then runs stats::model.matrix() on the result.
 - If, in the original mold() call, indicators = "none" was set, it runs stats::model.matrix() on the result without the factor columns, and then adds them on afterwards.
 - If, in the original mold() call, indicators = "one_hot" was set, it runs stats::model.matrix() on the result with a contrast function that includes indicators for all levels of all factor columns.
 - If any offsets are present from using offset() in the original call to mold(), then they are extracted with model_offset().
 - If intercept = TRUE in the original call to mold(), then an intercept column is added.
 - It coerces the result of the above steps to a tibble.
- Outcomes
 - It runs stats::model.frame() on new_data using the stored terms object corresponding to the outcomes.
 - Coerces the result to a tibble.

Differences From Base R

There are a number of differences from base R regarding how formulas are processed by mold() that require some explanation.

Multivariate outcomes can be specified on the LHS using syntax that is similar to the RHS (i.e. outcome_1 + outcome_2 ~ predictors). If any complex calculations are done on the LHS and they return matrices (like stats::poly()), then those matrices are flattened into multiple columns of the tibble after the call to model.frame(). While this is possible, it is not recommended, and if a large amount of preprocessing is required on the outcomes, then you are better off using a recipes::recipe().

Global variables are *not* allowed in the formula. An error will be thrown if they are included. All terms in the formula should come from data. If you need to use inline functions in the formula, the safest way to do so is to prefix them with their package name, like pkg::fn(). This ensures that the function will always be available at mold() (fit) and forge() (prediction) time. That said, if the package is *attached* (i.e. with library()), then you should be able to use the inline function without the prefix.

By default, intercepts are *not* included in the predictor output from the formula. To include an intercept, set blueprint = default_formula_blueprint(intercept = TRUE). The rationale for this is that many packages either always require or never allow an intercept (for example, the earth package), and they do a large amount of extra work to keep the user from supplying one or removing it. This interface standardizes all of that flexibility in one place.

```
# ------
data("hardhat-example-data")
# ------
# Formula Example
# Call mold() with the training data
processed <- mold(</pre>
 log(num_1) ~ num_2 + fac_1,
  example_train,
 blueprint = default_formula_blueprint(intercept = TRUE)
# Then, call forge() with the blueprint and the test data
# to have it preprocess the test data in the same way
forge(example_test, processed$blueprint)
# Use `outcomes = TRUE` to also extract the preprocessed outcome
forge(example_test, processed$blueprint, outcomes = TRUE)
# -----
# Factors without an intercept
# No intercept is added by default
processed <- mold(num_1 ~ fac_1 + fac_2, example_train)</pre>
\# So, for factor columns, the first factor is completely expanded into all
\mbox{\ensuremath{\mbox{\#}}}\ \mbox{\ensuremath{\mbox{`K'}}}\ \mbox{\ensuremath{\mbox{columns}}}\ \mbox{\ensuremath{\mbox{(the number of levels)}}\ , and the subsequent factors are expanded
# into `K - 1` columns.
processed$predictors
# In the above example, `fac_1` is expanded into all three columns,
# `fac_2` is not. This behavior comes from `model.matrix()`, and is somewhat
\mbox{\#} known in the R community, but can lead to a model that is difficult to
# interpret since the corresponding p-values are testing wildly different
# hypotheses.
# To get all indicators for all columns (irrespective of the intercept),
# use the `indicators = "one_hot"` option
processed <- mold(</pre>
 num_1 ~ fac_1 + fac_2,
  example_train,
 blueprint = default_formula_blueprint(indicators = "one_hot")
processed$predictors
# It is not possible to construct a no-intercept model that expands all
# factors into `K - 1` columns using the formula method. If required, a
# recipe could be used to construct this model.
# ------
# Global variables
```

```
y <- rep(1, times = nrow(example_train))</pre>
# In base R, global variables are allowed in a model formula
frame <- model.frame(fac_1 ~ y + num_2, example_train)</pre>
head(frame)
# mold() does not allow them, and throws an error
try(mold(fac_1 ~ y + num_2, example_train))
# -----
# Dummy variables and interactions
# By default, factor columns are expanded
# and interactions are created, both by
# calling `model.matrix()`. Some models (like
# tree based models) can take factors directly
# but still might want to use the formula method.
# In those cases, set `indicators = "none"` to not
# run `model.matrix()` on factor columns. Interactions
# are still allowed and are run on numeric columns.
bp_no_indicators <- default_formula_blueprint(indicators = "none")</pre>
processed <- mold(</pre>
  ~ fac_1 + num_1:num_2,
  example_train,
 blueprint = bp_no_indicators
processed$predictors
# An informative error is thrown when `indicators = "none"` and
# factors are present in interaction terms or in inline functions
try(mold(num_1 ~ num_2:fac_1, example_train, blueprint = bp_no_indicators))
try(mold(num_1 ~ paste0(fac_1), example_train, blueprint = bp_no_indicators))
# -----
# Multivariate outcomes
# Multivariate formulas can be specified easily
processed <- mold(num_1 + log(num_2) ~ fac_1, example_train)</pre>
processed$outcomes
# Inline functions on the LHS are run, but any matrix
# output is flattened (like what happens in `model.matrix()`)
# (essentially this means you don't wind up with columns
# in the tibble that are matrices)
processed <- mold(poly(num_2, degree = 2) ~ fac_1, example_train)</pre>
processed$outcomes
# TRUE
ncol(processed$outcomes) == 2
# Multivariate formulas specified in mold()
# carry over into forge()
forge(example_test, processed$blueprint, outcomes = TRUE)
```

```
# Offsets
# Offsets are handled specially in base R, so they deserve special
# treatment here as well. You can add offsets using the inline function
# `offset()`
processed <- mold(num_1 ~ offset(num_2) + fac_1, example_train)</pre>
processed$extras$offset
# Multiple offsets can be included, and they get added together
processed <- mold(</pre>
 num_1 ~ offset(num_2) + offset(num_3),
  example_train
)
identical(
  processed$extras$offset$.offset,
 example_train$num_2 + example_train$num_3
# Forging test data will also require
# and include the offset
forge(example_test, processed$blueprint)
# ------
# Intercept only
# Because '1' and '0' are intercept modifying terms, they are
# not allowed in the formula and are instead controlled by the
# `intercept` argument of the blueprint. To use an intercept
# only formula, you should supply `NULL` on the RHS of the formula.
mold(
  ~NULL,
  example_train,
 blueprint = default_formula_blueprint(intercept = TRUE)
# ------
# Matrix output for predictors
# You can change the `composition` of the predictor data set
bp <- default_formula_blueprint(composition = "dgCMatrix")</pre>
processed <- mold(log(num_1) ~ num_2 + fac_1, example_train, blueprint = bp)</pre>
class(processed$predictors)
```

default_recipe_blueprint

Default recipe blueprint

Description

This pages holds the details for the recipe preprocessing blueprint. This is the blueprint used by default from mold() if x is a recipe.

Usage

```
default_recipe_blueprint(
  intercept = FALSE,
  allow_novel_levels = FALSE,
  fresh = TRUE,
  bake_dependent_roles = character(),
  composition = "tibble"
)

## S3 method for class 'recipe'
mold(x, data, ..., blueprint = NULL)
```

Arguments

intercept

A logical. Should an intercept be included in the processed data? This information is used by the process function in the mold and forge function list.

allow_novel_levels

A logical. Should novel factor levels be allowed at prediction time? This information is used by the clean function in the forge function list, and is passed on to scream().

fresh Should already trained operations be re-trained when prep() is called? bake_dependent_roles

A character vector of recipes column "roles" specifying roles that are required to recipes::bake() new data. Can't be "predictor" or "outcome", as predictors are always required and outcomes are handled by the outcomes argument of forge().

Typically, non-standard roles (such as "id" or "case_weights") are not required to bake() new data. Unless specified by bake_dependent_roles, these non-standard role columns are excluded from checks done in forge() to validate the column structure of new_data, will not be passed to bake() even if they existed in new_data, and will not be returned in the forge()\$extras\$roles slot. See the documentation of recipes::add_role() for more information about roles.

composition

Either "tibble", "matrix", or "dgCMatrix" for the format of the processed predictors. If "matrix" or "dgCMatrix" are chosen, all of the predictors must be numeric after the preprocessing method has been applied; otherwise an error is thrown.

x An unprepped recipe created from recipes::recipe().

data A data frame or matrix containing the outcomes and predictors.

... Not used.

blueprint A preprocessing blueprint. If left as NULL, then a default_recipe_blueprint()

is used.

Value

For default_recipe_blueprint(), a recipe blueprint.

Mold

When mold() is used with the default recipe blueprint:

- It calls recipes::prep() to prep the recipe.
- It calls recipes::juice() to extract the outcomes and predictors. These are returned as tibbles
- If intercept = TRUE, adds an intercept column to the predictors.

Forge

When forge() is used with the default recipe blueprint:

- It calls shrink() to trim new_data to only the required columns and coerce new_data to a tibble.
- It calls scream() to perform validation on the structure of the columns of new_data.
- It calls recipes::bake() on the new_data using the prepped recipe used during training.
- It adds an intercept column onto new_data if intercept = TRUE.

```
library(recipes)
# ------
# Setup
train <- iris[1:100, ]</pre>
test <- iris[101:150, ]
# ------
# Recipes example
# Create a recipe that logs a predictor
rec <- recipe(Species ~ Sepal.Length + Sepal.Width, train) %>%
 step_log(Sepal.Length)
processed <- mold(rec, train)</pre>
# Sepal.Length has been logged
processed$predictors
processed$outcomes
# The underlying blueprint is a prepped recipe
processed$blueprint$recipe
# Call forge() with the blueprint and the test data
# to have it preprocess the test data in the same way
forge(test, processed$blueprint)
# Use 'outcomes = TRUE' to also extract the preprocessed outcome!
# This logged the Sepal.Length column of `new_data`
forge(test, processed$blueprint, outcomes = TRUE)
# -----
```

```
# With an intercept
# You can add an intercept with `intercept = TRUE`
processed <- mold(rec, train, blueprint = default_recipe_blueprint(intercept = TRUE))</pre>
processed$predictors
# But you also could have used a recipe step
rec2 <- step_intercept(rec)</pre>
mold(rec2, iris)$predictors
# ------
# Non standard roles
# If you have custom recipes roles, it is assumed that they are required
# in `prep()` and afterwards for modeling, but are not required at `bake()`
# time and for prediction. This means that they are processed and returned
# in the `$extras$roles` slot of the return value of `mold()`, but they
# are not required to be in `new_data` in `forge()` and are not returned
# in the result.
rec_roles <- recipe(train) %>%
  update_role(Sepal.Width, new_role = "predictor") %>%
  update_role(Species, new_role = "outcome") %>%
  update_role(Sepal.Length, new_role = "id") %>%
  update_role(Petal.Length, new_role = "important")
processed_roles <- mold(rec_roles, train)</pre>
# The custom roles will still be in the `mold()` result in case you need
# them for modeling.
processed_roles$extras
# Notice that the columns with custom roles exist in `test`,
# but they weren't passed to `bake()` and aren't in the output.
forge(test, processed_roles$blueprint)$extras
# They can even be removed from `test` entirely, and it still works.
test2 <- test
test2$Petal.Length <- NULL
forge(test2, processed_roles$blueprint)$extras
# Occasionally, you might have a custom role that is required to be able
# to `bake()` on `new_data`. In those cases, you can specify it with
# `bake_dependent_roles` in `default_recipe_blueprint()`, which will ensure
# that it is a required column when calling `forge()`, that it will be
# passed on to `bake()`, and that it will be returned in the result.
bp <- default_recipe_blueprint(bake_dependent_roles = "important")</pre>
processed_roles <- mold(rec_roles, train, blueprint = bp)</pre>
# Now `"important"` is a required role when `forge()`-ing
forge(test, processed_roles$blueprint)$extras$roles
# Which means that we can't `forge()` with the data frame that is missing
# the `Petal.Length` column
```

```
try(forge(test2, processed_roles$blueprint))
# ------
# Matrix output for predictors

# You can change the `composition` of the predictor data set
bp <- default_recipe_blueprint(composition = "dgCMatrix")
processed <- mold(rec, train, blueprint = bp)
class(processed$predictors)</pre>
```

 $default_xy_blueprint$ $Default\ XY\ blueprint$

Description

This pages holds the details for the XY preprocessing blueprint. This is the blueprint used by default from mold() if x and y are provided separately (i.e. the XY interface is used).

Usage

```
default_xy_blueprint(
  intercept = FALSE,
  allow_novel_levels = FALSE,
  composition = "tibble"
)
## S3 method for class 'data.frame'
mold(x, y, ..., blueprint = NULL)
## S3 method for class 'matrix'
mold(x, y, ..., blueprint = NULL)
```

Arguments

intercept

A logical. Should an intercept be included in the processed data? This information is used by the process function in the mold and forge function list

allow_novel_levels

A logical. Should novel factor levels be allowed at prediction time? This information is used by the clean function in the forge function list, and is passed on to scream().

composition

Either "tibble", "matrix", or "dgCMatrix" for the format of the processed predictors. If "matrix" or "dgCMatrix" are chosen, all of the predictors must be numeric after the preprocessing method has been applied; otherwise an error is thrown.

A data frame or matrix containing the predictors.

y A data frame, matrix, or vector containing the outcomes.

.. Not used.

blueprint A preprocessing blueprint. If left as NULL, then a default_xy_blueprint()

is used.

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Details

As documented in **standardize()**, if y is a *vector*, then the returned outcomes tibble has 1 column with a standardized name of ".outcome".

The one special thing about the XY method's forge function is the behavior of outcomes = TRUE when a *vector* y value was provided to the original call to mold(). In that case, mold() converts y into a tibble, with a default name of .outcome. This is the column that forge() will look for in new_data to preprocess. See the examples section for a demonstration of this.

Value

For default_xy_blueprint(), an XY blueprint.

Mold

When mold() is used with the default xy blueprint:

- It converts **x** to a tibble.
- It adds an intercept column to x if intercept = TRUE.
- It runs standardize() on y.

Forge

When forge() is used with the default xy blueprint:

- It calls shrink() to trim new_data to only the required columns and coerce new_data to a tibble.
- It calls scream() to perform validation on the structure of the columns of new_data.
- It adds an intercept column onto new_data if intercept = TRUE.

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```
# ------
# Intercept
processed <- mold(train_x, train_y, blueprint = default_xy_blueprint(intercept = TRUE))</pre>
forge(test_x, processed$blueprint)
# -----
# XY Method and forge(outcomes = TRUE)
# You can request that the new outcome columns are preprocessed as well, but
# they have to be present in `new_data`!
processed <- mold(train_x, train_y)</pre>
# Can't do this!
try(forge(test_x, processed$blueprint, outcomes = TRUE))
\mbox{\#} 
 Need to use the full test set, including 'y'
forge(test, processed$blueprint, outcomes = TRUE)
# With the XY method, if the Y value used in `mold()` is a vector,
# then a column name of `.outcome` is automatically generated.
# This name is what forge() looks for in `new_data`.
# Y is a vector!
y_vec <- train_y$Species</pre>
processed_vec <- mold(train_x, y_vec)</pre>
# This throws an informative error that tell you
# to include an `".outcome"` column in `new_data`.
try(forge(iris, processed_vec$blueprint, outcomes = TRUE))
test2 <- test
test2$.outcome <- test2$Species</pre>
test2$Species <- NULL
# This works, and returns a tibble in the $outcomes slot
forge(test2, processed_vec$blueprint, outcomes = TRUE)
# -----
# Matrix output for predictors
# You can change the `composition` of the predictor data set
bp <- default_xy_blueprint(composition = "dgCMatrix")</pre>
processed <- mold(train_x, train_y, blueprint = bp)</pre>
class(processed$predictors)
```

delete_response

Delete the response from a terms object

Description

delete_response() is exactly the same as delete.response(), except that it fixes a long

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standing bug by also removing the part of the "dataClasses" attribute corresponding to the response, if it exists.

Usage

```
delete_response(terms)
```

Arguments

terms

A terms object.

Details

The bug is described here:

```
https://stat.ethz.ch/pipermail/r-devel/2012-January/062942.html
```

Value

terms with the response sections removed.

Examples

```
framed <- model_frame(Species ~ Sepal.Width, iris)
attr(delete.response(framed$terms), "dataClasses")
attr(delete_response(framed$terms), "dataClasses")</pre>
```

forge

Forge prediction-ready data

Description

forge() applies the transformations requested by the specific blueprint on a set of new_data. This new_data contains new predictors (and potentially outcomes) that will be used to generate predictions.

All blueprints have consistent return values with the others, but each is unique enough to have its own help page. Click through below to learn how to use each one in conjunction with forge().

- XY Method default_xy_blueprint()
- Formula Method default_formula_blueprint()
- Recipes Method default_recipe_blueprint()

Usage

```
forge(new_data, blueprint, ..., outcomes = FALSE)
```

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Arguments

new_data A data frame or matrix of predictors to process. If outcomes = TRUE, this

should also contain the outcomes to process.

blueprint A preprocessing blueprint.

... Not used.

outcomes A logical. Should the outcomes be processed and returned as well?

Details

If the outcomes are present in <code>new_data</code>, they can optionally be processed and returned in the <code>outcomes</code> slot of the returned list by setting <code>outcomes = TRUE</code>. This is very useful when doing cross validation where you need to preprocess the outcomes of a test set before computing performance.

Value

A named list with 3 elements:

- predictors: A tibble containing the preprocessed new_data predictors.
- outcomes: If outcomes = TRUE, a tibble containing the preprocessed outcomes found in new_data. Otherwise, NULL.
- extras: Either NULL if the blueprint returns no extra information, or a named list containing the extra information.

Examples

```
# See the blueprint specific documentation linked above
# for various ways to call forge with different
# blueprints.

train <- iris[1:100, ]
test <- iris[101:150, ]

# Formula
processed <- mold(
   log(Sepal.Width) ~ Species,
   train,
   blueprint = default_formula_blueprint(indicators = "none")
)

forge(test, processed$blueprint, outcomes = TRUE)</pre>
```

 $frequency_weights$

Frequency weights

Description

[Experimental]

frequency_weights() creates a vector of frequency weights which allow you to compactly repeat an observation a set number of times. Frequency weights are supplied as a non-negative integer vector, where only whole numbers are allowed.

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Usage

```
frequency_weights(x)
```

Arguments

Χ

An integer vector.

Value

A new frequency weights vector.

See Also

```
importance_weights()
```

Examples

```
# Record that the first observation has 10 replicates, the second has 12
# replicates, and so on
frequency_weights(c(10, 12, 2, 1))
# Fractional values are not allowed
try(frequency_weights(c(1.5, 2.3, 10)))
```

get_data_classes

Extract data classes from a data frame or matrix

Description

When predicting from a model, it is often important for the new_data to have the same classes as the original data used to fit the model. get_data_classes() extracts the classes from the original training data.

Usage

```
get_data_classes(data)
```

Arguments

data

A data frame or matrix.

Value

A named list. The names are the column names of data and the values are character vectors containing the class of that column.

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Examples

```
get_data_classes(iris)
get_data_classes(as.matrix(mtcars))
# Unlike .MFclass(), the full class
# vector is returned
data <- data.frame(col = ordered(c("a", "b")))
.MFclass(data$col)
get_data_classes(data)</pre>
```

get_levels

Extract factor levels from a data frame

Description

get_levels() extracts the levels from any factor columns in data. It is mainly useful for extracting the original factor levels from the predictors in the training set. get_outcome_levels()
is a small wrapper around get_levels() for extracting levels from a factor outcome that
first calls standardize() on y.

Usage

```
get_levels(data)
get_outcome_levels(y)
```

Arguments

data

A data frame to extract levels from.

У

The outcome. This can be:

- A factor vector
- A numeric vector
- A 1D numeric array
- A numeric matrix with column names
- A 2D numeric array with column names
- A data frame with numeric or factor columns

Value

A named list with as many elements as there are factor columns in data or y. The names are the names of the factor columns, and the values are character vectors of the levels.

If there are no factor columns, NULL is returned.

See Also

```
stats::.getXlevels()
```

Examples

```
# Factor columns are returned with their levels
get_levels(iris)

# No factor columns
get_levels(mtcars)

# standardize() is first run on `y`
# which converts the input to a data frame
# with an automatically named column, `".outcome"`
get_outcome_levels(y = factor(letters[1:5]))
```

hardhat-example-data Example data for hardhat

Description

Example data for hardhat

Details

Data objects for a training and test set with the same variables: three numeric and two factor columns.

Value

Examples

```
data("hardhat-example-data")
```

hardhat-extract

Generics for object extraction

Description

These generics are used to extract elements from various model objects. Methods are defined in other packages, such as tune, workflows, and workflowsets, but the returned object is always the same.

- extract_fit_engine() returns the engine specific fit embedded within a parsnip model fit. For example, when using parsnip::linear_reg() with the "lm" engine, this returns the underlying lm object.
- extract_fit_parsnip() returns a parsnip model fit.
- extract_mold() returns the preprocessed "mold" object returned from mold(). It contains information about the preprocessing, including either the prepped recipe, the formula terms object, or variable selectors.

- extract_spec_parsnip() returns a parsnip model specification.
- extract_preprocessor() returns the formula, recipe, or variable expressions used for preprocessing.
- extract_recipe() returns a recipe, possibly estimated.
- extract_workflow() returns a workflow, possibly fit.
- extract_parameter_dials() returns a single dials parameter object.
- extract_parameter_set_dials() returns a set of dials parameter objects.

Usage

```
extract_workflow(x, ...)
extract_recipe(x, ...)
extract_spec_parsnip(x, ...)
extract_fit_parsnip(x, ...)
extract_fit_engine(x, ...)
extract_mold(x, ...)
extract_preprocessor(x, ...)
extract_parameter_dials(x, ...)
extract_parameter_set_dials(x, ...)
```

Arguments

x An object.

... Extra arguments passed on to methods.

Examples

```
\# See packages where methods are defined for examples, such as 'parsnip' or \# 'workflows'.
```

importance_weights

Importance weights

Description

[Experimental]

importance_weights() creates a vector of importance weights which allow you to apply a context dependent weight to your observations. Importance weights are supplied as a non-negative double vector, where fractional values are allowed.

Usage

```
importance_weights(x)
```

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Arguments

x A double vector.

Value

A new importance weights vector.

See Also

```
frequency_weights()
```

Examples

```
importance_weights(c(1.5, 2.3, 10))
```

 $is_blueprint$

Is x a preprocessing blueprint?

Description

```
is_blueprint() checks if x inherits from "hardhat_blueprint".
```

Usage

```
is_blueprint(x)
```

Arguments

x

An object.

Examples

```
is_blueprint(default_xy_blueprint())
```

is_case_weights

Is x a case weights vector?

${\bf Description}$

[Experimental]

is_case_weights() checks if x inherits from "hardhat_case_weights".

Usage

```
is_case_weights(x)
```

Arguments

Χ

An object.

Value

A single TRUE or FALSE.

Examples

```
is_case_weights(1)
is_case_weights(frequency_weights(1))
```

is_frequency_weights

Is \times a frequency weights vector?

Description

[Experimental]

 $is_frequency_weights()\ checks\ if\ x\ inherits\ from\ "hardhat_frequency_weights".$

Usage

```
is_frequency_weights(x)
```

Arguments

Χ

An object.

Value

A single TRUE or FALSE.

Examples

```
is_frequency_weights(1)
is_frequency_weights(frequency_weights(1))
is_frequency_weights(importance_weights(1))
```

is_importance_weights Is x an importance weights vector?

Description

[Experimental]

is_importance_weights() checks if x inherits from "hardhat_importance_weights".

Usage

```
is_importance_weights(x)
```

Arguments

Χ

An object.

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Value

A single TRUE or FALSE.

Examples

```
is_importance_weights(1)
is_importance_weights(frequency_weights(1))
is_importance_weights(importance_weights(1))
```

model_frame

Construct a model frame

Description

model_frame() is a stricter version of stats::model.frame(). There are a number of differences, with the main being that rows are *never* dropped and the return value is a list with the frame and terms separated into two distinct objects.

Usage

```
model_frame(formula, data)
```

Arguments

formula A formula or terms object representing the terms of the model frame.

data A data frame or matrix containing the terms of formula.

Details

The following explains the rationale for some of the difference in arguments compared to stats::model.frame():

- subset: Not allowed because the number of rows before and after model_frame() has been run should always be the same.
- na.action: Not allowed and is forced to "na.pass" because the number of rows before and after model_frame() has been run should always be the same.
- drop.unused.levels: Not allowed because it seems inconsistent for data and the result
 of model_frame() to ever have the same factor column but with different levels, unless
 specified though original_levels. If this is required, it should be done through a
 recipe step explicitly.
- xlev: Not allowed because this check should have been done ahead of time. Use scream() to check the integrity of data against a training set if that is required.
- ...: Not exposed because offsets are handled separately, and it is not necessary to pass weights here any more because rows are never dropped (so weights don't have to be subset alongside the rest of the design matrix). If other non-predictor columns are required, use the "roles" features of recipes.

It is important to always use the results of model_frame() with model_matrix() rather than stats::model.matrix() because the tibble in the result of model_frame() does not have a terms object attached. If model.matrix(;terms;, ;tibble;) is called directly, then a call to model.frame() will be made automatically, which can give faulty results.

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Value

A named list with two elements:

- "data": A tibble containing the model frame.
- "terms": A terms object containing the terms for the model frame.

Examples

model_matrix

Construct a design matrix

Description

model_matrix() is a stricter version of stats::model.matrix(). Notably, model_matrix() will never drop rows, and the result will be a tibble.

Usage

```
model_matrix(terms, data)
```

Arguments

terms A terms object to construct a model matrix with. This is typically the

terms object returned from the corresponding call to model_frame().

data A tibble to construct the design matrix with. This is typically the tibble

returned from the corresponding call to model_frame().

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Details

The following explains the rationale for some of the difference in arguments compared to stats::model.matrix():

- contrasts.arg: Set the contrasts argument, options("contrasts") globally, or assign a contrast to the factor of interest directly using stats::contrasts(). See the examples section.
- xlev: Not allowed because model.frame() is never called, so it is unnecessary.
- ...: Not allowed because the default method of model.matrix() does not use it, and the lm method uses it to pass potential offsets and weights through, which are handled differently in hardhat.

Value

A tibble containing the design matrix.

```
# ------
# Example usage
framed <- model_frame(Sepal.Width ~ Species, iris)</pre>
model_matrix(framed$terms, framed$data)
# ------
# Missing values never result in dropped rows
iris2 <- iris</pre>
iris2$Species[1] <- NA</pre>
framed2 <- model_frame(Sepal.Width ~ Species, iris2)</pre>
model_matrix(framed2$terms, framed2$data)
# -----
# Contrasts
# Default contrasts
y <- factor(c("a", "b"))</pre>
x < - data.frame(y = y)
framed <- model_frame(~y, x)</pre>
# Setting contrasts directly
y_with_contrast <- y</pre>
contrasts(y_with_contrast) <- contr.sum(2)</pre>
x2 <- data.frame(y = y_with_contrast)
framed2 <- model_frame(~y, x2)</pre>
# Compare!
model_matrix(framed$terms, framed$data)
model_matrix(framed2$terms, framed2$data)
# Also, can set the contrasts globally
global_override <- c(unordered = "contr.sum", ordered = "contr.poly")</pre>
```

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```
rlang::with_options(
    .expr = {
      model_matrix(framed$terms, framed$data)
    },
    contrasts = global_override
)
```

model_offset

Extract a model offset

Description

model_offset() extracts a numeric offset from a model frame. It is inspired by stats::model.offset(), but has nicer error messages and is slightly stricter.

Usage

```
model_offset(terms, data)
```

Arguments

terms A "terms" object corresponding to data, returned from a call to model_frame().

data A data frame returned from a call to model_frame().

Details

If a column that has been tagged as an offset is not numeric, a nice error message is thrown telling you exactly which column was problematic.

stats::model.offset() also allows for a column named "(offset)" to be considered an offset along with any others that have been tagged by stats::offset(). However, stats::model.matrix()
does not recognize these columns as offsets (so it doesn't remove them as it should).
Because of this inconsistency, columns named "(offset)" are not treated specially by
model_offset().

Value

A numeric vector representing the offset.

```
x <- model.frame(Species ~ offset(Sepal.Width), iris)
model_offset(terms(x), x)

xx <- model.frame(Species ~ offset(Sepal.Width) + offset(Sepal.Length), iris)
model_offset(terms(xx), xx)

# Problematic columns are caught with intuitive errors
tryCatch(
    expr = {</pre>
```

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```
x <- model.frame(~ offset(Species), iris)
  model_offset(terms(x), x)
},
error = function(e) {
  print(e$message)
}
</pre>
```

mold

Mold data for modeling

Description

mold() applies the appropriate processing steps required to get training data ready to be fed into a model. It does this through the use of various *blueprints* that understand how to preprocess data that come in various forms, such as a formula or a recipe.

All blueprints have consistent return values with the others, but each is unique enough to have its own help page. Click through below to learn how to use each one in conjunction with mold().

- XY Method default_xy_blueprint()
- Formula Method default_formula_blueprint()
- Recipes Method default_recipe_blueprint()

Usage

```
mold(x, ...)
```

Arguments

x An object. See the method specific implementations linked in the Description for more information.

... Not used.

Value

A named list containing 4 elements:

- predictors: A tibble containing the molded predictors to be used in the model.
- outcome: A tibble containing the molded outcomes to be used in the model.
- blueprint: A method specific "hardhat_blueprint" object for use when making predictions.
- extras: Either NULL if the blueprint returns no extra information, or a named list containing the extra information.

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Examples

```
# See the method specific documentation linked in Description
# for the details of each blueprint, and more examples.

# XY
mold(iris[, "Sepal.Width", drop = FALSE], iris$Species)

# Formula
mold(Species ~ Sepal.Width, iris)

# Recipe
library(recipes)
mold(recipe(Species ~ Sepal.Width, iris), iris)
```

new_case_weights

Extend case weights

Description

[Experimental]

new_case_weights() is a developer oriented function for constructing a new case weights type. The ¡case_weights¿ type itself is an *abstract* type with very little functionality. Because of this, class is a required argument.

Usage

```
new_case_weights(x, ..., class)
```

Arguments

x An integer or double vector.

... Name-value pairs defining attributes

class Name of subclass.

Value

A new subclassed case weights vector.

```
new_case_weights(1:5, class = "my_weights")
```

new_default_formula_blueprint

Create a new default blueprint

Description

This page contains the constructors for the default blueprints. They can be extended if you want to add extra behavior on top of what the default blueprints already do, but generally you will extend the non-default versions of the constructors found in the documentation for new_blueprint().

Usage

```
new_default_formula_blueprint(
  mold,
  forge,
  intercept = FALSE,
  allow_novel_levels = FALSE,
  ptypes = NULL,
  formula = NULL,
  indicators = "traditional",
  composition = "tibble",
  terms = list(predictors = NULL, outcomes = NULL),
  subclass = character()
)
new_default_recipe_blueprint(
  mold,
  forge,
  intercept = FALSE,
  allow_novel_levels = FALSE,
  fresh = TRUE,
  bake_dependent_roles = character(),
  composition = "tibble",
  ptypes = NULL,
  recipe = NULL,
  extra_role_ptypes = NULL,
  subclass = character()
)
new_default_xy_blueprint(
  mold,
  forge,
  intercept = FALSE,
  allow_novel_levels = FALSE,
  composition = "tibble",
  ptypes = NULL,
  subclass = character()
```

)

Arguments

mold A named list with two elements, clean and process, see the new_blueprint()

section, Mold Functions, for details.

forge A named list with two elements, clean and process, see the new_blueprint()

section, Forge Functions, for details.

intercept A logical. Should an intercept be included in the processed data? This in-

> formation is used by the process function in the mold and forge function list.

allow_novel_levels

A logical. Should novel factor levels be allowed at prediction time? This information is used by the clean function in the forge function list, and

is passed on to scream().

ptypes Either NULL, or a named list with 2 elements, predictors and outcomes,

> both of which are 0-row tibbles. ptypes is generated automatically at mold() time and is used to validate new_data at prediction time. At mold() time, the information found in blueprint\$mold\$process()\$ptype

is used to set ptypes for the blueprint.

Either NULL, or a formula that specifies how the predictors and outcomes

should be preprocessed. This argument is set automatically at mold()

time.

A single character string. Control how factors are expanded into dummy variable indicator columns. One of:

> • "traditional" - The default. Create dummy variables using the traditional model.matrix() infrastructure. Generally this creates K -1 indicator columns for each factor, where K is the number of levels in that factor.

"none" - Leave factor variables alone. No expansion is done.

"one_hot" - Create dummy variables using a one-hot encoding approach that expands unordered factors into all K indicator columns,

rather than K -1.

Either "tibble", "matrix", or "dgCMatrix" for the format of the processed composition

predictors. If "matrix" or "dgCMatrix" are chosen, all of the predictors must be numeric after the preprocessing method has been applied; oth-

erwise an error is thrown.

terms A named list of two elements, predictors and outcomes. Both elements are terms objects that describe the terms for the outcomes and predictors

separately. This argument is set automatically at mold() time.

Name-value pairs for additional elements of blueprints that subclass this

blueprint.

subclass A character vector. The subclasses of this blueprint.

Should already trained operations be re-trained when prep() is called? fresh

bake_dependent_roles

A character vector of recipes column "roles" specifying roles that are required to recipes::bake() new data. Can't be "predictor" or "outcome", as predictors are always required and outcomes are handled by the outcomes argument of forge().

formula

indicators

Typically, non-standard roles (such as "id" or "case_weights") are not required to bake() new data. Unless specified by bake_dependent_roles, these non-standard role columns are excluded from checks done in forge() to validate the column structure of new_data, will not be passed to bake() even if they existed in new_data, and will not be returned in the forge()\$extras\$roles slot. See the documentation of recipes::add_role() for more information about roles.

recipe

Either NULL, or an unprepped recipe. This argument is set automatically at mold() time.

extra_role_ptypes

A named list. The names are the unique non-standard recipe roles (i.e. everything except "predictors" and "outcomes"). The values are prototypes of the original columns with that role. These are used for validation in forge().

new_formula_blueprint Create a new preprocessing blueprint

Description

These are the base classes for creating new preprocessing blueprints. All blueprints inherit from the one created by new_blueprint(), and the default method specific blueprints inherit from the other three here.

If you want to create your own processing blueprint for a specific method, generally you will subclass one of the method specific blueprints here. If you want to create a completely new preprocessing blueprint for a totally new preprocessing method (i.e. not the formula, xy, or recipe method) then you should subclass new_blueprint().

Usage

```
new_formula_blueprint(
  mold,
  forge,
  intercept = FALSE,
  allow_novel_levels = FALSE,
  ptypes = NULL,
  formula = NULL,
  indicators = "traditional",
  composition = "tibble",
  subclass = character()
)
new_recipe_blueprint(
  mold,
  forge,
  intercept = FALSE,
  allow_novel_levels = FALSE,
  fresh = TRUE,
  bake_dependent_roles = character(),
```

```
composition = "tibble",
  ptypes = NULL,
  recipe = NULL,
  subclass = character()
new_xy_blueprint(
  mold,
  forge,
  intercept = FALSE,
  allow_novel_levels = FALSE,
  composition = "tibble",
  ptypes = NULL,
  subclass = character()
new_blueprint(
  mold,
  forge,
  intercept = FALSE,
  allow_novel_levels = FALSE,
  composition = "tibble",
  ptypes = NULL,
  subclass = character()
)
```

Arguments

mold A named list with two elements, clean and process, see the new_blueprint()

section, Mold Functions, for details.

forge A named list with two elements, clean and process, see the new_blueprint()

section, Forge Functions, for details.

intercept A logical. Should an intercept be included in the processed data? This in-

formation is used by the process function in the mold and forge function

list.

allow_novel_levels

A logical. Should novel factor levels be allowed at prediction time? This information is used by the clean function in the forge function list, and

is passed on to scream().

ptypes Either NULL, or a named list with 2 elements, predictors and outcomes,

both of which are 0-row tibbles. ptypes is generated automatically at mold() time and is used to validate new_data at prediction time. At mold() time, the information found in blueprint\$mold\$process()\$ptype

is used to set ptypes for the blueprint.

formula Either NULL, or a formula that specifies how the predictors and outcomes

should be preprocessed. This argument is set automatically at mold()

time.

indicators A single character string. Control how factors are expanded into dummy

variable indicator columns. One of:

- "traditional" The default. Create dummy variables using the traditional model.matrix() infrastructure. Generally this creates K
 -1 indicator columns for each factor, where K is the number of levels in that factor.
- "none" Leave factor variables alone. No expansion is done.
- "one_hot" Create dummy variables using a one-hot encoding approach that expands unordered factors into all K indicator columns, rather than K -1.

composition

Either "tibble", "matrix", or "dgCMatrix" for the format of the processed predictors. If "matrix" or "dgCMatrix" are chosen, all of the predictors must be numeric after the preprocessing method has been applied; otherwise an error is thrown.

.. Name-value pairs for additional elements of blueprints that subclass this blueprint.

subclass A character vector. The subclasses of this blueprint.

fresh Should already trained operations be re-trained when prep() is called? bake_dependent_roles

A character vector of recipes column "roles" specifying roles that are required to recipes::bake() new data. Can't be "predictor" or "outcome", as predictors are always required and outcomes are handled by the outcomes argument of forge().

Typically, non-standard roles (such as "id" or "case_weights") are not required to bake() new data. Unless specified by bake_dependent_roles, these non-standard role columns are excluded from checks done in forge() to validate the column structure of new_data, will not be passed to bake() even if they existed in new_data, and will not be returned in the forge()\$extras\$roles slot. See the documentation of recipes::add_role() for more information about roles.

recipe

Either NULL, or an unprepped recipe. This argument is set automatically at mold() time.

Value

A preprocessing blueprint, which is a list containing the inputs used as arguments to the function, along with a class specific to the type of blueprint being created.

Mold Functions

blueprint\$mold should be a named list with two elements, both of which are functions:

- clean: A function that performs initial cleaning of the user's input data to be used in the model.
 - Arguments:
 - * If this is an xy blueprint, blueprint, x and y.
 - * Otherwise, blueprint and data.
 - Output: A named list of three elements:
 - * blueprint: The blueprint, returned and potentially updated.
 - * If using an xy blueprint:
 - · x: The cleaned predictor data.
 - · y: The cleaned outcome data.

- * If not using an xy blueprint:
 - data: The cleaned data.
- process: A function that performs the actual preprocessing of the data.
 - Arguments:
 - * If this is an xy blueprint, blueprint, x and y.
 - * Otherwise, blueprint and data.
 - Output: A named list of 5 elements:
 - * blueprint: The blueprint, returned and potentially updated.
 - * predictors: A tibble of predictors.
 - * outcomes: A tibble of outcomes.
 - * ptypes: A named list with 2 elements, predictors and outcomes, where both elements are 0-row tibbles.
 - * extras: Varies based on the blueprint. If the blueprint has no extra information, NULL. Otherwise a named list of the extra elements returned by the blueprint.

Both blueprint\$mold\$clean() and blueprint\$mold\$process() will be called, in order, from mold().

Forge Functions

blueprint\$forge should be a named list with two elements, both of which are functions:

- clean: A function that performs initial cleaning of new_data:
 - Arguments:
 - * blueprint, new_data, and outcomes.
 - Output: A named list of the following elements:
 - * blueprint: The blueprint, returned and potentially updated.
 - * predictors: A tibble containing the cleaned predictors.
 - * outcomes: A tibble containing the cleaned outcomes.
 - * extras: A named list of any extras obtained while cleaning. These are passed on to the process() function for further use.
- process: A function that performs the actual preprocessing of the data using the known information in the blueprint.
 - Arguments:
 - * blueprint, new_data, outcomes, extras.
 - Output: A named list of the following elements:
 - * blueprint: The blueprint, returned and potentially updated.
 - * predictors: A tibble of the predictors.
 - * outcomes: A tibble of the outcomes, or NULL.
 - * extras: Varies based on the blueprint. If the blueprint has no extra information, NULL. Otherwise a named list of the extra elements returned by the blueprint.

Both blueprint\$forge\$clean() and blueprint\$forge\$process() will be called, in order, from forge().

new_frequency_weights Construct a frequency weights vector

Description

[Experimental]

new_frequency_weights() is a developer oriented function for constructing a new frequency weights vector. Generally, you should use frequency_weights() instead.

Usage

```
new_frequency_weights(x = integer(), ..., class = character())
```

Arguments

x An integer vector.

... Name-value pairs defining attributes

class Name of subclass.

Value

A new frequency weights vector.

Examples

```
new_frequency_weights()
new_frequency_weights(1:5)
```

new_importance_weights Construct an importance weights vector

Description

[Experimental]

new_importance_weights() is a developer oriented function for constructing a new importance weights vector. Generally, you should use importance_weights() instead.

Usage

```
new_importance_weights(x = double(), ..., class = character())
```

Arguments

x A double vector.

... Name-value pairs defining attributes

class Name of subclass.

Value

A new importance weights vector.

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Examples

```
new_importance_weights()
new_importance_weights(c(1.5, 2.3, 10))
```

new_model

Constructor for a base model

Description

A model is a *scalar object*, as classified in Advanced R. As such, it takes uniquely named elements in ... and combines them into a list with a class of class. This entire object represent a single model.

Usage

```
new_model(..., blueprint = default_xy_blueprint(), class = character())
```

Arguments

... Name-value pairs for elements specific to the model defined by class.

blueprint A preprocessing blueprint returned from a call to mold().

class A character vector representing the class of the model.

Details

Because every model should have multiple interfaces, including formula and recipes interfaces, all models should have a blueprint that can process new data when predict() is called. The easiest way to generate an blueprint with all of the information required at prediction time is to use the one that is returned from a call to mold().

Value

A new scalar model object, represented as a classed list with named elements specified in

```
new_model(
  custom_element = "my-elem",
  blueprint = default_xy_blueprint(),
  class = "custom_model"
)
```

refresh_blueprint 39

refresh_blueprint

Refresh a preprocessing blueprint

Description

refresh_blueprint() is a developer facing generic function that is called at the end of update_blueprint(). It simply is a wrapper around the method specific new_*_blueprint() function that runs the updated blueprint through the constructor again to ensure that all of the elements of the blueprint are still valid after the update.

Usage

```
refresh_blueprint(blueprint)
```

Arguments

blueprint

A preprocessing blueprint.

Details

If you implement your own custom blueprint, you should export a refresh_blueprint() method that just calls the constructor for your blueprint and passes through all of the elements of the blueprint to the constructor.

Value

blueprint is returned after a call to the corresponding constructor.

```
blueprint <- default_xy_blueprint()

# This should never be done manually, but is essentially
# what `update_blueprint(blueprint, intercept = TRUE)` does for you
blueprint$intercept <- TRUE

# Then update_blueprint() will call refresh_blueprint()
# to ensure that the structure is correct
refresh_blueprint(blueprint)

# So you can't do something like...
blueprint_bad <- blueprint
blueprint_bad$intercept <- 1

# ...because the constructor will catch it
try(refresh_blueprint(blueprint_bad))

# And update_blueprint() catches this automatically
try(update_blueprint(blueprint, intercept = 1))</pre>
```

40 scream

run mold

Call mold\$clean() and mold\$process()

Description

This is a purely developer facing function, that is *only* used if you are creating a completely new blueprint inheriting only from new_blueprint(), and not from one of the more common: new_xy_blueprint(), new_recipe_blueprint(), new_formula_blueprint().

Usage

```
run_mold(blueprint, ...)
```

Arguments

blueprint A preprocessing blueprint.

... Not used. Required for extensibility.

Details

Because mold() has different interfaces (like XY and formula), which require different arguments (x and y vs data), their corresponding blueprints also have different arguments for the blueprint\$mold\$clean() and blueprint\$mold\$process() functions. The sole job of run_mold() is simply to call these two functions with the right arguments.

The only time you need to implement a method for run_mold() is if you are creating a new_blueprint() that does not follow one of the three core blueprint types. In that special case, create a method for run_mold() with your blueprint type, and pass through whatever arguments are necessary to call your blueprint specific clean() and process() functions.

If you go this route, you will also need to create a mold() method if x is not a data frame / matrix, recipe, or formula. If x is one of those types, then run_mold() will be called for you by the existing mold() method, you just have to supply the run_mold() method for your blueprint.

Value

The preprocessed result, as a named list.

scream Scream.

Description

scream() ensures that the structure of data is the same as prototype, ptype. Under the
hood, vctrs::vec_cast() is used, which casts each column of data to the same type as the
corresponding column in ptype.

This casting enforces a number of important structural checks, including but not limited to:

• Data Classes - Checks that the class of each column in data is the same as the corresponding column in ptype.

scream 41

• Novel Levels - Checks that the factor columns in data don't have any new levels when compared with the ptype columns. If there are new levels, a warning is issued and they are coerced to NA. This check is optional, and can be turned off with allow_novel_levels = TRUE.

• Level Recovery - Checks that the factor columns in data aren't missing any factor levels when compared with the ptype columns. If there are missing levels, then they are restored.

Usage

scream(data, ptype, allow_novel_levels = FALSE)

Arguments

data

A data frame containing the new data to check the structure of.

ptype

A data frame prototype to cast data to. This is commonly a 0-row slice of the training set.

allow_novel_levels

Should novel factor levels in data be allowed? The safest approach is the default, which throws a warning when novel levels are found, and coerces them to NA values. Setting this argument to TRUE will ignore all novel levels. This argument does not apply to ordered factors. Novel levels are not allowed in ordered factors because the level ordering is a critical part of the type.

Details

scream() is called by forge() after shrink() but before the actual processing is done. Generally, you don't need to call scream() directly, as forge() will do it for you.

If scream() is used as a standalone function, it is good practice to call shrink() right before it as there are no checks in scream() that ensure that all of the required column names actually exist in data. Those checks exist in shrink().

Value

A tibble containing the required columns after any required structural modifications have been made.

Factor Levels

scream() tries to be helpful by recovering missing factor levels and warning about novel levels. The following graphic outlines how scream() handles factor levels when coercing from a column in data to a column in ptype.

То	Factor			Ordered				
From	Same levels	New levels (allowed)	New levels (prohibited)	Missing levels	Same levels	New levels (allowed)	New levels (prohibited)	Missing levels
Factor	No change	No change	Warning: New levels coerced to NA	Levels recovered	Error	Error	Error	Error
Ordered	Error	Error	Error	Error	No change	Error	Error	Error
Character	Coerce to factor	Coerce to factor	Coerce to factor + Warning: New levels coerced to NA	Coerce to factor + Levels recovered	Coerce to ordered	Error	Error	Coerce to ordered + Levels recovered

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Note that ordered factor handing is much stricter than factor handling. Ordered factors in data must have *exactly* the same levels as ordered factors in ptype.

```
______
# Setup
train <- iris[1:100, ]</pre>
test <- iris[101:150, ]
# mold() is run at model fit time
# and a formula preprocessing blueprint is recorded
x <- mold(log(Sepal.Width) ~ Species, train)</pre>
# Inside the result of mold() are the prototype tibbles
# for the predictors and the outcomes
ptype_pred <- x$blueprint$ptypes$predictors</pre>
ptype_out <- x$blueprint$ptypes$outcomes</pre>
# shrink() / scream()
# Pass the test data, along with a prototype, to
# shrink() to extract the prototype columns
test_shrunk <- shrink(test, ptype_pred)</pre>
# Now pass that to scream() to perform validation checks
# If no warnings / errors are thrown, the checks were
# successful!
scream(test_shrunk, ptype_pred)
# ------
# Outcomes
# To also extract the outcomes, use the outcome prototype
test_outcome <- shrink(test, ptype_out)</pre>
scream(test_outcome, ptype_out)
# Casting
# scream() uses vctrs::vec_cast() to intelligently convert
# new data to the prototype automatically. This means
# it can automatically perform certain conversions, like
# coercing character columns to factors.
test2 <- test
test2$Species <- as.character(test2$Species)</pre>
test2_shrunk <- shrink(test2, ptype_pred)</pre>
scream(test2_shrunk, ptype_pred)
# It can also recover missing factor levels.
# For example, it is plausible that the test data only had the
# "virginica" level
test3 <- test
test3$Species <- factor(test3$Species, levels = "virginica")</pre>
```

shrink 43

```
test3_shrunk <- shrink(test3, ptype_pred)</pre>
test3_fixed <- scream(test3_shrunk, ptype_pred)</pre>
# scream() recovered the missing levels
levels(test3_fixed$Species)
# Novel levels
# When novel levels with any data are present in `data`, the default
# is to coerce them to `NA` values with a warning.
test4 <- test
test4$Species <- as.character(test4$Species)</pre>
test4$Species[1] <- "new_level"</pre>
test4$Species <- factor(</pre>
  test4$Species,
 levels = c(levels(test$Species), "new_level")
test4 <- shrink(test4, ptype_pred)</pre>
# Warning is thrown
test4_removed <- scream(test4, ptype_pred)</pre>
# Novel level is removed
levels(test4_removed$Species)
# No warning is thrown
test4_kept <- scream(test4, ptype_pred, allow_novel_levels = TRUE)</pre>
# Novel level is kept
levels(test4_kept$Species)
```

shrink

Subset only required columns

Description

shrink() subsets data to only contain the required columns specified by the prototype,
ptype.

Usage

```
shrink(data, ptype)
```

Arguments

data A data frame containing the data to subset.

ptype A data frame prototype containing the required columns.

Details

shrink() is called by forge() before scream() and before the actual processing is done.

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Value

A tibble containing the required columns.

Examples

```
# Setup
train <- iris[1:100, ]</pre>
test <- iris[101:150, ]
# -----
# shrink()
# mold() is run at model fit time
# and a formula preprocessing blueprint is recorded
x <- mold(log(Sepal.Width) ~ Species, train)</pre>
# Inside the result of mold() are the prototype tibbles
# for the predictors and the outcomes
ptype_pred <- x$blueprint$ptypes$predictors</pre>
ptype_out <- x$blueprint$ptypes$outcomes</pre>
# Pass the test data, along with a prototype, to
# shrink() to extract the prototype columns
shrink(test, ptype_pred)
# To extract the outcomes, just use the
# outcome prototype
shrink(test, ptype_out)
# shrink() makes sure that the columns
# required by `ptype` actually exist in the data
# and errors nicely when they don't
test2 <- subset(test, select = -Species)</pre>
try(shrink(test2, ptype_pred))
```

spruce

Spruce up predictions

Description

The family of spruce_*() functions convert predictions into a standardized format. They are generally called from a prediction implementation function for the specific type of prediction to return.

Usage

```
spruce_numeric(pred)
spruce_class(pred_class)
spruce_prob(pred_levels, prob_matrix)
```

standardize 45

Arguments

- pred_levels should be a character vector of the original levels of the outcome used in training.
- prob_matrix should be a numeric matrix of class probabilities with as many columns as levels in pred_levels, and in the same order.

Details

After running a spruce_*() function, you should *always* use the validation function validate_prediction_size() to ensure that the number of rows being returned is the same as the number of rows in the input (new_data).

Value

A tibble, ideally with the same number of rows as the new_data passed to predict(). The column names and number of columns vary based on the function used, but are standardized.

standardize

Standardize the outcome

Description

Most of the time, the input to a model should be flexible enough to capture a number of different input types from the user. standardize() focuses on capturing the flexibility in the *outcome*.

Usage

```
standardize(y)
```

Arguments

У

The outcome. This can be:

- A factor vector
- A numeric vector
- A 1D numeric array
- A numeric matrix with column names
- A 2D numeric array with column names
- A data frame with numeric or factor columns

Details

standardize() is called from mold() when using an XY interface (i.e. a y argument was supplied).

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Value

All possible values of y are transformed into a tibble for standardization. Vectors are transformed into a tibble with a single column named ".outcome".

Examples

```
standardize(1:5)
standardize(factor(letters[1:5]))
mat <- matrix(1:10, ncol = 2)
colnames(mat) <- c("a", "b")
standardize(mat)

df <- data.frame(x = 1:5, y = 6:10)
standardize(df)</pre>
```

tune

Mark arguments for tuning

Description

tune() is an argument placeholder to be used with the recipes, parsnip, and tune packages. It marks recipes step and parsnip model arguments for tuning.

Usage

```
tune(id = "")
```

Arguments

id

A single character value that can be used to differentiate parameters that are used in multiple places but have the same name, or if the user wants to add a note to the specified parameter.

Value

A call object that echos the user's input.

See Also

```
tune::tune_grid(), tune::tune_bayes()
```

```
tune()
tune("your name here")

# In practice, `tune()` is used alongside recipes or parsnip to mark
# specific arguments for tuning
library(recipes)

recipe(mpg ~ ., data = mtcars) %>%
    step_normalize(all_numeric_predictors()) %>%
    step_pca(all_numeric_predictors, num_comp = tune())
```

update_blueprint 47

Update a preprocessing blueprint	update_blueprint
----------------------------------	------------------

Description

update_blueprint() is the correct way to alter elements of an existing blueprint object. It has two benefits over just doing blueprint\$elem <-new_elem.

- The name you are updating *must* already exist in the blueprint. This prevents you from accidentally updating non-existent elements.
- The constructor for the blueprint is automatically run after the update by refresh_blueprint() to ensure that the blueprint is still valid.

Usage

```
update_blueprint(blueprint, ...)
```

Arguments

blueprint A preprocessing blueprint.

... Name-value pairs of *existing* elements in blueprint that should be updated.

Examples

```
blueprint <- default_xy_blueprint()

# `intercept` defaults to FALSE
blueprint

update_blueprint(blueprint, intercept = TRUE)

# Can't update non-existent elements
try(update_blueprint(blueprint, intercpt = TRUE))

# Can't add non-valid elements
try(update_blueprint(blueprint, intercept = 1))</pre>
```

 $ext{validate_column_names}$ $ext{\it Ensure that data contains required column names}$

Description

validate - asserts the following:

• The column names of data must contain all original_names.

check - returns the following:

- ok A logical. Does the check pass?
- missing_names A character vector. The missing column names.

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Usage

```
validate_column_names(data, original_names)
check_column_names(data, original_names)
```

Arguments

```
data A data frame to check.

original_names A character vector. The original column names.
```

Details

A special error is thrown if the missing column is named ".outcome". This only happens in the case where mold() is called using the xy-method, and a *vector* y value is supplied rather than a data frame or matrix. In that case, y is coerced to a data frame, and the automatic name ".outcome" is added, and this is what is looked for in forge(). If this happens, and the user tries to request outcomes using forge(...,outcomes = TRUE) but the supplied new_data does not contain the required ".outcome" column, a special error is thrown telling them what to do. See the examples!

Value

```
validate_column_names() returns data invisibly.
check_column_names() returns a named list of two components, ok, and missing_names.
```

Validation

hardhat provides validation functions at two levels.

- check_*(): check a condition, and return a list. The list always contains at least one element, ok, a logical that specifies if the check passed. Each check also has check specific elements in the returned list that can be used to construct meaningful error messages.
- validate_*(): check a condition, and error if it does not pass. These functions call their corresponding check function, and then provide a default error message. If you, as a developer, want a different error message, then call the check_*() function yourself, and provide your own validation function.

See Also

```
Other validation functions: validate_no_formula_duplication(), validate_outcomes_are_binary(), validate_outcomes_are_factors(), validate_outcomes_are_numeric(), validate_outcomes_are_univariate() validate_prediction_size(), validate_predictors_are_numeric()
```

```
foriginal_names <- colnames(mtcars)

test <- mtcars
bad_test <- test[, -c(3, 4)]

# All good</pre>
```

```
check_column_names(test, original_names)
# Missing 2 columns
check_column_names(bad_test, original_names)
# Will error
try(validate_column_names(bad_test, original_names))
# -----
# Special error when `.outcome` is missing
train <- iris[1:100, ]</pre>
test <- iris[101:150, ]
train_x <- subset(train, select = -Species)</pre>
train_y <- train$Species</pre>
# Here, y is a vector
processed <- mold(train_x, train_y)</pre>
# So the default column name is `".outcome"`
processed$outcomes
# It doesn't affect forge() normally
forge(test, processed$blueprint)
# But if the outcome is requested, and `".outcome"`
# is not present in `new_data`, an error is thrown
# with very specific instructions
try(forge(test, processed$blueprint, outcomes = TRUE))
# To get this to work, just create an .outcome column in new_data
test$.outcome <- test$Species</pre>
forge(test, processed$blueprint, outcomes = TRUE)
```

validate_no_formula_duplication

Ensure no duplicate terms appear in formula

Description

validate - asserts the following:

- formula must not have duplicates terms on the left and right hand side of the formula.
- check returns the following:
 - ok A logical. Does the check pass?
 - duplicates A character vector. The duplicate terms.

Usage

```
validate_no_formula_duplication(formula, original = FALSE)
check_no_formula_duplication(formula, original = FALSE)
```

Arguments

formula A formula to check.

original A logical. Should the original names be checked, or should the names

after processing be used? If FALSE, y $^{\sim}$ log(y) is allowed because the names are "y" and "log(y)", if TRUE, y $^{\sim}$ log(y) is not allowed because

the original names are both "y".

Value

validate_no_formula_duplication() returns formula invisibly.
check_no_formula_duplication() returns a named list of two components, ok and duplicates.

Validation

hardhat provides validation functions at two levels.

- check_*(): check a condition, and return a list. The list always contains at least one
 element, ok, a logical that specifies if the check passed. Each check also has check
 specific elements in the returned list that can be used to construct meaningful error
 messages.
- validate_*(): check a condition, and error if it does not pass. These functions call their corresponding check function, and then provide a default error message. If you, as a developer, want a different error message, then call the check_*() function yourself, and provide your own validation function.

See Also

Other validation functions: validate_column_names(), validate_outcomes_are_binary(), validate_outcomes_are_factors(), validate_outcomes_are_numeric(), validate_outcomes_are_univariate() validate_prediction_size(), validate_predictors_are_numeric()

```
# All good
check_no_formula_duplication(y ~ x)

# Not good!
check_no_formula_duplication(y ~ y)

# This is generally okay
check_no_formula_duplication(y ~ log(y))

# But you can be more strict
check_no_formula_duplication(y ~ log(y), original = TRUE)

# This would throw an error
try(validate_no_formula_duplication(log(y) ~ log(y)))
```

validate_outcomes_are_binary

Ensure that the outcome has binary factors

Description

validate - asserts the following:

• outcomes must have binary factor columns.

check - returns the following:

- ok A logical. Does the check pass?
- bad_cols A character vector. The names of the columns with problems.
- num_levels An integer vector. The actual number of levels of the columns with problems.

Usage

```
validate_outcomes_are_binary(outcomes)
check_outcomes_are_binary(outcomes)
```

Arguments

outcomes

An object to check.

Details

The expected way to use this validation function is to supply it the \$outcomes element of the result of a call to mold().

Value

validate_outcomes_are_binary() returns outcomes invisibly.

 $\label{limits} {\tt check_outcomes_are_binary()} \ \ {\tt returns} \ \ {\tt a} \ \ {\tt named} \ \ {\tt list} \ \ {\tt of} \ \ {\tt three} \ \ {\tt components}, \ {\tt ok}, \ {\tt bad_cols}, \ {\tt and} \ \ {\tt num_levels}.$

Validation

hardhat provides validation functions at two levels.

- check_*(): check a condition, and return a list. The list always contains at least one element, ok, a logical that specifies if the check passed. Each check also has check specific elements in the returned list that can be used to construct meaningful error messages.
- validate_*(): check a condition, and error if it does not pass. These functions call their corresponding check function, and then provide a default error message. If you, as a developer, want a different error message, then call the check_*() function yourself, and provide your own validation function.

See Also

Other validation functions: validate_column_names(), validate_no_formula_duplication(), validate_outcomes_are_factors(), validate_outcomes_are_numeric(), validate_outcomes_are_univariate() validate_prediction_size(), validate_predictors_are_numeric()

Examples

```
# Not a binary factor. 0 levels
check_outcomes_are_binary(data.frame(x = 1))
# Not a binary factor. 1 level
check_outcomes_are_binary(data.frame(x = factor("A")))
# All good
check_outcomes_are_binary(data.frame(x = factor(c("A", "B"))))
```

validate_outcomes_are_factors

Ensure that the outcome has only factor columns

Description

validate - asserts the following:

• outcomes must have factor columns.

check - returns the following:

- ok A logical. Does the check pass?
- bad_classes A named list. The names are the names of problematic columns, and the values are the classes of the matching column.

Usage

```
validate_outcomes_are_factors(outcomes)
check_outcomes_are_factors(outcomes)
```

Arguments

outcomes An object to check.

Details

The expected way to use this validation function is to supply it the \$outcomes element of the result of a call to mold().

Value

```
validate_outcomes_are_factors() returns outcomes invisibly. check_outcomes_are_factors() returns a named list of two components, ok and bad_classes.
```

Validation

hardhat provides validation functions at two levels.

- check_*(): check a condition, and return a list. The list always contains at least one
 element, ok, a logical that specifies if the check passed. Each check also has check
 specific elements in the returned list that can be used to construct meaningful error
 messages.
- validate_*(): check a condition, and error if it does not pass. These functions call their corresponding check function, and then provide a default error message. If you, as a developer, want a different error message, then call the check_*() function yourself, and provide your own validation function.

See Also

```
Other validation functions: validate_column_names(), validate_no_formula_duplication(), validate_outcomes_are_binary(), validate_outcomes_are_numeric(), validate_outcomes_are_univariate(), validate_prediction_size(), validate_predictors_are_numeric()
```

Examples

```
# Not a factor column.
check_outcomes_are_factors(data.frame(x = 1))
# All good
check_outcomes_are_factors(data.frame(x = factor(c("A", "B"))))
```

validate_outcomes_are_numeric

Ensure outcomes are all numeric

Description

validate - asserts the following:

• outcomes must have numeric columns.

check - returns the following:

- ok A logical. Does the check pass?
- bad_classes A named list. The names are the names of problematic columns, and the values are the classes of the matching column.

Usage

```
validate_outcomes_are_numeric(outcomes)
check_outcomes_are_numeric(outcomes)
```

Arguments

outcomes An object to check.

Details

The expected way to use this validation function is to supply it the \$outcomes element of the result of a call to mold().

Value

validate_outcomes_are_numeric() returns outcomes invisibly. check_outcomes_are_numeric() returns a named list of two components, ok and bad_classes.

Validation

hardhat provides validation functions at two levels.

- check_*(): check a condition, and return a list. The list always contains at least one element, ok, a logical that specifies if the check passed. Each check also has check specific elements in the returned list that can be used to construct meaningful error messages.
- validate_*(): check a condition, and error if it does not pass. These functions call their corresponding check function, and then provide a default error message. If you, as a developer, want a different error message, then call the check_*() function yourself, and provide your own validation function.

See Also

Other validation functions: validate_column_names(), validate_no_formula_duplication(), validate_outcomes_are_binary(), validate_outcomes_are_factors(), validate_outcomes_are_univariate(), validate_prediction_size(), validate_predictors_are_numeric()

Examples

```
# All good
check_outcomes_are_numeric(mtcars)

# Species is not numeric
check_outcomes_are_numeric(iris)

# This gives an intelligent error message
try(validate_outcomes_are_numeric(iris))
```

validate_outcomes_are_univariate

Ensure that the outcome is univariate

Description

validate - asserts the following:

• outcomes must have 1 column. Atomic vectors are treated as 1 column matrices.

check - returns the following:

- ok A logical. Does the check pass?
- n_cols A single numeric. The actual number of columns.

Usage

```
validate_outcomes_are_univariate(outcomes)
check_outcomes_are_univariate(outcomes)
```

Arguments

outcomes

An object to check.

Details

The expected way to use this validation function is to supply it the \$outcomes element of the result of a call to mold().

Value

```
validate_outcomes_are_univariate() returns outcomes invisibly.

check_outcomes_are_univariate() returns a named list of two components, ok and n_cols.
```

Validation

hardhat provides validation functions at two levels.

- check_*(): check a condition, and return a list. The list always contains at least one element, ok, a logical that specifies if the check passed. Each check also has check specific elements in the returned list that can be used to construct meaningful error messages.
- validate_*(): check a condition, and error if it does not pass. These functions call their corresponding check function, and then provide a default error message. If you, as a developer, want a different error message, then call the check_*() function yourself, and provide your own validation function.

See Also

```
Other validation functions: validate_column_names(), validate_no_formula_duplication(), validate_outcomes_are_binary(), validate_outcomes_are_factors(), validate_outcomes_are_numeric(), validate_prediction_size(), validate_predictors_are_numeric()
```

```
validate_outcomes_are_univariate(data.frame(x = 1))
try(validate_outcomes_are_univariate(mtcars))
```

validate_prediction_size

Ensure that predictions have the correct number of rows

Description

validate - asserts the following:

• The size of pred must be the same as the size of new_data.

check - returns the following:

- ok A logical. Does the check pass?
- size_new_data A single numeric. The size of new_data.
- size_pred A single numeric. The size of pred.

Usage

```
validate_prediction_size(pred, new_data)
check_prediction_size(pred, new_data)
```

Arguments

pred A tibble. The predictions to return from any prediction type. This is

often created using one of the spruce functions, like spruce_numeric().

new_data A data frame of new predictors and possibly outcomes.

Details

This validation function is one that is more developer focused rather than user focused. It is a final check to be used right before a value is returned from your specific predict() method, and is mainly a "good practice" sanity check to ensure that your prediction blueprint always returns the same number of rows as new_data, which is one of the modeling conventions this package tries to promote.

Value

validate_prediction_size() returns pred invisibly.

check_prediction_size() returns a named list of three components, ok, size_new_data, and size_pred.

Validation

hardhat provides validation functions at two levels.

- check_*(): check a condition, and return a list. The list always contains at least one element, ok, a logical that specifies if the check passed. Each check also has check specific elements in the returned list that can be used to construct meaningful error messages.
- validate_*(): check a condition, and error if it does not pass. These functions call their corresponding check function, and then provide a default error message. If you, as a developer, want a different error message, then call the check_*() function yourself, and provide your own validation function.

See Also

Other validation functions: validate_column_names(), validate_no_formula_duplication(), validate_outcomes_are_binary(), validate_outcomes_are_factors(), validate_outcomes_are_numeric(), validate_outcomes_are_univariate(), validate_predictors_are_numeric()

Examples

```
# Say new_data has 5 rows
new_data <- mtcars[1:5, ]

# And somehow you generate predictions
# for those 5 rows
pred_vec <- 1:5

# Then you use `spruce_numeric()` to clean
# up these numeric predictions
pred <- spruce_numeric(pred_vec)

pred

# Use this check to ensure that
# the number of rows or pred match new_data
check_prediction_size(pred, new_data)

# An informative error message is thrown
# if the rows are different
try(validate_prediction_size(spruce_numeric(1:4), new_data))</pre>
```

 $validate_predictors_are_numeric$

Ensure predictors are all numeric

Description

validate - asserts the following:

• predictors must have numeric columns.

check - returns the following:

- ok A logical. Does the check pass?
- bad_classes A named list. The names are the names of problematic columns, and the values are the classes of the matching column.

Usage

```
validate_predictors_are_numeric(predictors)
check_predictors_are_numeric(predictors)
```

Arguments

predictors An object to check.

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Details

The expected way to use this validation function is to supply it the \$predictors element of the result of a call to mold().

Value

validate_predictors_are_numeric() returns predictors invisibly.
check_predictors_are_numeric() returns a named list of two components, ok, and bad_classes.

Validation

hardhat provides validation functions at two levels.

- check_*(): check a condition, and return a list. The list always contains at least one
 element, ok, a logical that specifies if the check passed. Each check also has check
 specific elements in the returned list that can be used to construct meaningful error
 messages.
- validate_*(): check a condition, and error if it does not pass. These functions call their corresponding check function, and then provide a default error message. If you, as a developer, want a different error message, then call the check_*() function yourself, and provide your own validation function.

See Also

Other validation functions: validate_column_names(), validate_no_formula_duplication(), validate_outcomes_are_binary(), validate_outcomes_are_factors(), validate_outcomes_are_numeric(), validate_outcomes_are_univariate(), validate_prediction_size()

Examples

```
# All good
check_predictors_are_numeric(mtcars)
# Species is not numeric
check_predictors_are_numeric(iris)
# This gives an intelligent error message
try(validate_predictors_are_numeric(iris))
```

weighted_table

 $Weighted\ table$

Description

weighted_table() computes a weighted contingency table based on factors provided in ... and a double vector of weights provided in weights. It can be seen as a weighted extension to base::table() and an alternative to stats::xtabs().

weighted_table() always uses the *exact* set of levels returned by levels() when constructing the table. This results in the following properties:

Missing values found in the factors are never included in the table unless there is an
explicit NA factor level. If needed, this can be added to a factor with base::addNA()
or forcats::fct_expand(x,NA).

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• Levels found in the factors that aren't actually used in the underlying data are included in the table with a value of 0. If needed, you can drop unused factor levels by re-running your factor through factor(), or by calling forcats::fct_drop().

See the examples section for more information about these properties.

Usage

```
weighted_table(..., weights, na_remove = FALSE)
```

Arguments

	Factors of equal length to use in the weighted table. If the are named, those names will propagate onto the "dimnames names" of the resulting table. At least one factor must be provided.
weights	A double vector of weights used to fill the cells of the weighted table. This must be the same length as the factors provided in
na_remove	A single TRUE or FALSE for handling whether or not missing values in weights should be removed when summing up the weights.

Details

The result of weighted_table() does not have a "table" class attached to it. It is only a double array. This is because "table" objects are defined as containing integer counts, but weighted tables can utilize fractional weights.

Value

The weighted table as an array of double values.

```
x <- factor(c("x", "y", "z", "x", "x", "y"))
y <- factor(c("a", "b", "a", "a", "b", "b"))</pre>
w \leftarrow c(1.5, 2, 1.1, .5, 3, 2)
weighted_table(x = x, y = y, weights = w)
# If `weights` contains missing values, then missing values will be
# propagated into the weighted table
x <- factor(c("x", "y", "y"))</pre>
y <- factor(c("a", "b", "b"))
w < -c(1, NA, 3)
weighted_table(x = x, y = y, weights = w)
# You can remove the missing values while summing up the weights with
# `na_remove = TRUE`
weighted_table(x = x, y = y, weights = w, na_remove = TRUE)
# ------
# If there are missing values in the factors, those typically don't show
# up in the weighted table
x <- factor(c("x", NA, "y", "x"))
y <- factor(c("a", "b", "a", NA))
```

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```
w <- 1:4
weighted_table(x = x, y = y, weights = w)
# This is because the missing values aren't considered explicit levels
levels(x)
# You can force them to show up in the table by using `addNA()` ahead of time
# (or `forcats::fct_expand(x, NA)`)
x \leftarrow addNA(x, ifany = TRUE)
y \leftarrow addNA(y, ifany = TRUE)
levels(x)
weighted_table(x = x, y = y, weights = w)
# If there are levels in your factors that aren't actually used in the
\mbox{\#} underlying data, then they will still show up in the table with a '0' value
x <- factor(c("x", "y", "x"), levels = c("x", "y", "z"))
y <- factor(c("a", "b", "a"), levels = c("a", "b", "c"))</pre>
w <- 1:3
weighted_table(x = x, y = y, weights = w)
# If you want to drop these empty factor levels from the result, you can
# rerun `factor()` ahead of time to drop them (or `forcats::fct_drop()`)
x <- factor(x)</pre>
y <- factor(y)</pre>
levels(x)
weighted_table(x = x, y = y, weights = w)
```

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