Package ‘rsample’

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Title General Resampling Infrastructure
Version 0.0.7
Maintainer Max Kuhn <max@rstudio.com>
Description Classes and functions to create and summarize different types of resampling objects (e.g. bootstrap, cross-validation).
Imports dplyr (>= 0.8.5), purrr, tibble, rlang (>= 0.4.0), methods, generics, utils, tidyselect, furrr, tidyr, vctrs (>= 0.3.0)
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Author Max Kuhn [aut, cre],
Fanny Chow [aut],
Hadley Wickham [aut],
RStudio [cph]
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Description
For a data set, `add_resample_id()` will add at least one new column that identifies which resample that the data came from. In most cases, a single column is added but for some resampling methods two or more are added.

Usage
```
add_resample_id(.data, split, dots = FALSE)
```
apparent

Arguments

.data   A data frame
.split  A single rset object.
dots    A single logical: should the id columns be prefixed with a "." to avoid name conflicts with .data?

Value

An updated data frame.

See Also

labels.rsplit

Examples

library(dplyr)

set.seed(363)
car_folds <- vfold_cv(mtcars, repeats = 3)

analysis(car_folds$splits[[1]]) %>%
  add_resample_id(car_folds$splits[[1]]) %>%
  head()

car_bt <- bootstraps(mtcars)

analysis(car_bt$splits[[1]]) %>%
  add_resample_id(car_bt$splits[[1]]) %>%
  head()

---

Sampling for the Apparent Error Rate

Description

When building a model on a data set and re-predicting the same data, the performance estimate from those predictions is often call the “apparent” performance of the model. This estimate can be wildly optimistic. "Apparent sampling" here means that the analysis and assessment samples are the same. These resamples are sometimes used in the analysis of bootstrap samples and should otherwise be avoided like old sushi.

Usage

apparent(data, ...)
Arguments

data A data frame.
...

Value

A tibble with a single row and classes apparent, rset, tbl_df, tbl, and data.frame. The results include a column for the data split objects and one column called id that has a character string with the resample identifier.

Examples

apparent(mtcars)

as.data.frame.rsplit Convert an rsplit object to a data frame

Description

The analysis or assessment code can be returned as a data frame (as dictated by the data argument) using as.data.frame.rsplit. analysis and assessment are shortcuts.

Usage

## S3 method for class 'rsplit'
as.data.frame(x, row.names = NULL, optional = FALSE, data = "analysis", ...)

analysis(x, ...)

assessment(x, ...)

Arguments

x An rsplit object.
row.names NULL or a character vector giving the row names for the data frame. Missing values are not allowed.
optional A logical: should the column names of the data be checked for legality?
data Either "analysis" or "assessment" to specify which data are returned.
...

Examples

library(dplyr)
set.seed(104)
folds <- vfold_cv(mtcars)

model_data_1 <- folds$splits[[1]] %>% analysis()
holdout_data_1 <- folds$splits[[1]] %>% assessment()
attrition  

**Job Attrition**

**Description**
Job Attrition

**Details**
These data are from the IBM Watson Analytics Lab. The website describes the data with “Uncover the factors that lead to employee attrition and explore important questions such as ‘show me a breakdown of distance from home by job role and attrition’ or ‘compare average monthly income by education and attrition’. This is a fictional data set created by IBM data scientists.”. There are 1470 rows.

These data are now in the modeldata package.

---

bootstraps  

**Bootstrap Sampling**

**Description**
A bootstrap sample is a sample that is the same size as the original data set that is made using replacement. This results in analysis samples that have multiple replicates of some of the original rows of the data. The assessment set is defined as the rows of the original data that were not included in the bootstrap sample. This is often referred to as the "out-of-bag" (OOB) sample.

**Usage**
`bootstraps(data, times = 25, strata = NULL, breaks = 4, apparent = FALSE, ...)`

**Arguments**
- `data` A data frame.
- `times` The number of bootstrap samples.
- `strata` A variable that is used to conduct stratified sampling. When not `NULL`, each bootstrap sample is created within the stratification variable. This could be a single character value or a variable name that corresponds to a variable that exists in the data frame.
- `breaks` A single number giving the number of bins desired to stratify a numeric stratification variable.
- `apparent` A logical. Should an extra resample be added where the analysis and holdout subset are the entire data set. This is required for some estimators used by the `summary` function that require the apparent error rate.
- `...` Not currently used.
Details

The argument `apparent` enables the option of an additional "resample" where the analysis and assessment data sets are the same as the original data set. This can be required for some types of analysis of the bootstrap results. The `strata` argument is based on a similar argument in the random forest package where the bootstrap samples are conducted within the stratification variable. This can help ensure that the number of data points in the bootstrap sample is equivalent to the proportions in the original data set. (Strata below 10% of the total are pooled together.)

Value

An tibble with classes bootstraps, rset, tbl_df, tbl, and data.frame. The results include a column for the data split objects and a column called `id` that has a character string with the resample identifier.

Examples

```r
bootstraps(mtcars, times = 2)
bootstraps(mtcars, times = 2, apparent = TRUE)

library(purrr)
iris2 <- iris[1:130, ]

set.seed(13)
resample1 <- bootstraps(iris2, times = 3)
map_dbl(resample1$splits,
    function(x) {
        dat <- as.data.frame(x)$Species
        mean(dat == "virginica")
    })

set.seed(13)
resample2 <- bootstraps(iris2, strata = "Species", times = 3)
map_dbl(resample2$splits,
    function(x) {
        dat <- as.data.frame(x)$Species
        mean(dat == "virginica")
    })

set.seed(13)
resample3 <- bootstraps(iris2, strata = "Sepal.Length", breaks = 6, times = 3)
map_dbl(resample3$splits,
    function(x) {
        dat <- as.data.frame(x)$Species
        mean(dat == "virginica")
    })
```

---

#### Determine the Assessment Samples

| runoff | Determine the Assessment Samples |
drinks

Description

Given an rsplit object, complement will determine which of the data rows are contained in the assessment set. To save space, many of the rset objects will not contain indicies for the assessment split.

Usage

complement(x, ...)

Arguments

x An rsplit object
...
Not currently used

Value

A integer vector.

See Also

populate()

Examples

set.seed(28432)
fold_rs <- vfold_cv(mtcars)
head(fold_rs$splits[[1]]$in_id)
fold_rs$splits[[1]]$out_id
complement(fold_rs$splits[[1]])

drinks  Sample Time Series Data

Description

Sample Time Series Data

Details

Drink sales. The exact name of the series from FRED is: "Merchant Wholesalers, Except Manufacturers’ Sales Branches and Offices Sales: Nondurable Goods: Beer, Wine, and Distilled Alcoholic Beverages Sales" These data are now in the modeldata package.
form_pred

Extract Predictor Names from Formula or Terms

Description

call.vars returns all variables used in a formula. This function only returns the variables explicitly used on the right-hand side (i.e., it will not resolve dots unless the object is terms with a data set specified).

Usage

form_pred(object, ...)

Arguments

object A model formula or stats::terms() object.

... Arguments to pass to all.vars()

Value

A character vector of names

Examples

form_pred(y ~ x + z)
form_pred(terms(y ~ x + z))

form_pred(y ~ x + log(z))
form_pred(log(y) ~ x + z)

form_pred(y1 + y2 ~ x + z)
form_pred(log(y1) + y2 ~ x + z)

# will fail:
# form_pred(y ~ .)

form_pred(terms(Species ~ (.)^2, data = iris))
form_pred(terms(~ (.)^2, data = iris))
gather.rset

Gather an rset Object

Description

This method uses gather on an rset object to stack all of the non-ID or split columns in the data and is useful for stacking model evaluation statistics. The resulting data frame has a column based on the column names of data and another for the values.

Usage

```r
## S3 method for class 'rset'
gather(
  data,
  key = NULL,
  value = NULL,
  ..., 
  na.rm = TRUE,
  convert = FALSE,
  factor_key = TRUE
)
```

Arguments

data An rset object.

key, value, ... Not specified in this method and will be ignored. Note that this means that selectors are ignored if they are passed to the function.

na.rm If TRUE, will remove rows from output where the value column in NA.

convert If TRUE will automatically run type.convert() on the key column. This is useful if the column names are actually numeric, integer, or logical.

factor_key If FALSE, the default, the key values will be stored as a character vector. If TRUE, will be stored as a factor, which preserves the original ordering of the columns.

Value

A data frame with the ID columns, a column called `model` (with the previous column names), and a column called `statistic` (with the values).

Examples

```r
library(rsample)
cv_obj <- vfold_cv(mtcars, v = 10)
cv_obj$lm_rmse <- rnorm(10, mean = 2)
cv_obj$nnet_rmse <- rnorm(10, mean = 1)
gather(cv_obj)
```
**group_vfold_cv**

**Group V-Fold Cross-Validation**

**Description**

Group V-fold cross-validation creates splits of the data based on some grouping variable (which may have more than a single row associated with it). The function can create as many splits as there are unique values of the grouping variable or it can create a smaller set of splits where more than one value is left out at a time.

**Usage**

```r
group_vfold_cv(data, group = NULL, v = NULL, ...)
```

**Arguments**

- `data`: A data frame.
- `group`: This could be a single character value or a variable name that corresponds to a variable that exists in the data frame.
- `v`: The number of partitions of the data set. If let `NULL`, `v` will be set to the number of unique values in the group.
- `...`: Not currently used.

**Value**

An `tibble` with classes `group_vfold_cv`, `rset`, `tbl_df`, `tbl`, and `data.frame`. The results include a column for the data split objects and an identification variable.

**Examples**

```r
set.seed(3527)
test_data <- data.frame(id = sort(sample(1:20, size = 80, replace = TRUE)))
test_data$dat <- runif(nrow(test_data))

set.seed(5144)
split_by_id <- group_vfold_cv(test_data, group = "id")

get_id_left_out <- function(x)
  unique(assessment(x)$id)

library(purrr)
table(map_int(split_by_id$splits, get_id_left_out))

set.seed(5144)
split_by_some_id <- group_vfold_cv(test_data, group = "id", v = 7)
held_out <- map(split_by_some_id$splits, get_id_left_out)
table(unlist(held_out))
# number held out per resample:
map_int(held_out, length)
```
**initial_split**

*Simple Training/Test Set Splitting*

**Description**

`initial_split` creates a single binary split of the data into a training set and testing set. `initial_time_split` does the same, but takes the first prop samples for training, instead of a random selection. `training` and `testing` are used to extract the resulting data.

**Usage**

```r
initial_split(data, prop = 3/4, strata = NULL, breaks = 4, ...)
initial_time_split(data, prop = 3/4, lag = 0, ...)
training(x)
testing(x)
```

**Arguments**

- `data` A data frame.
- `prop` The proportion of data to be retained for modeling/analysis.
- `strata` A variable that is used to conduct stratified sampling to create the resamples. This could be a single character value or a variable name that corresponds to a variable that exists in the data frame.
- `breaks` A single number giving the number of bins desired to stratify a numeric stratification variable.
- `lag` A value to include an lag between the assessment and analysis set. This is useful if lagged predictors will be used during training and testing.
- `x` An `rsplit` object produced by `initial_split`

**Details**

The `strata` argument causes the random sampling to be conducted *within the stratification variable*. This can help ensure that the number of data points in the training data is equivalent to the proportions in the original data set. (Strata below 10% of the total are pooled together.)

**Value**

An `rsplit` object that can be used with the `training` and `testing` functions to extract the data in each split.
int_pctl

Bootstrap confidence intervals

Description

Calculate bootstrap confidence intervals using various methods.

Usage

int_pctl(.data, statistics, alpha = 0.05)

int_t(.data, statistics, alpha = 0.05)

int_bca(.data, statistics, alpha = 0.05, .fn, ...)

Arguments

.data A data frame containing the bootstrap resamples created using bootstraps(). For t- and BCa-intervals, the apparent argument should be set to TRUE. Even if the apparent argument is set to TRUE for the percentile method, the apparent data is never used in calculating the percentile confidence interval.

statistics An unquoted column name or dplyr selector that identifies a single column in the data set that contains the indiviual bootstrap estimates. This can be a list column of tidy tibbles (that contains columns term and estimate) or a simple numeric column. For t-intervals, a standard tidy column (usually called std.err) is required. See the examples below.

alpha Level of significance

.fn A function to calculate statistic of interest. The function should take an rsplit as the first argument and the ... are required.

... Arguments to pass to .fn.
Details

Percentile intervals are the standard method of obtaining confidence intervals but require thousands of resamples to be accurate. T-intervals may need fewer resamples but require a corresponding variance estimate. Bias-corrected and accelerated intervals require the original function that was used to create the statistics of interest and are computationally taxing.

Value

Each function returns a tibble with columns .lower, .estimate, .upper, .alpha, .method, and term. .method is the type of interval (eg. "percentile", "student-t", or "BCa"). term is the name of the estimate. Note the .estimate returned from int_pctl() is the mean of the estimates from the bootstrap resamples and not the estimate from the apparent model.

References

https://rsample.tidymodels.org/articles/Applications/Intervals.html

Examples

library(broom)
library(dplyr)
library(purrr)
library(tibble)

lm_est <- function(split, ...) {
  lm(mpg ~ disp + hp, data = analysis(split)) %>%
    tidy()
}

set.seed(52156)
car_rs <-
  bootstraps(mtcars, 500, apparent = TRUE) %>%
    mutate(results = map(splits, lm_est))

int_pctl(car_rs, results)
int_t(car_rs, results)
int_bca(car_rs, results, .fn = lm_est)

# putting results into a tidy format
rank_corr <- function(split) {
  dat <- analysis(split)
  tibble(
    term = "corr",
    estimate = cor(dat$Sepal.Length, dat$Sepal.Width, method = "spearman"),
    # don't know the analytical std.err so no t-intervals
    std.err = NA_real_
  )
}

}
```r
set.seed(69325)
bootstraps(iris, 500, apparent = TRUE) %>%
  mutate(correlations = map(splits, rank_corr)) %>%
  int_pctl(correlations)
```

---

**labels.rset**  
*Find Labels from rset Object*

**Description**  
Produce a vector of resampling labels (e.g. "Fold1") from an rset object. Currently, nested_cv is not supported.

**Usage**  
```r
## S3 method for class 'rset'
labels(object, make_factor = FALSE, ...)
```  
```r
## S3 method for class 'vfold_cv'
labels(object, make_factor = FALSE, ...)
```

**Arguments**  
- `object`: An rset object  
- `make_factor`: A logical for whether the results should be character or a factor.  
- `...`: Not currently used.

**Value**  
A single character or factor vector.

**Examples**  
```r
labels(vfold_cv(mtcars))
```

---

**labels.rsplit**  
*Find Labels from rsplit Object*

**Description**  
Produce a tibble of identification variables so that single splits can be linked to a particular resample.

**Usage**  
```r
## S3 method for class 'rsplit'
labels(object, ...)
```
Arguments

object  An rsplit object

Value

A tibble.

See Also

add_resample_id

Examples

```
cv_splits <- vfold_cv(mtcars)
labels(cv_splits$splits[[1]])
```

Description

Leave-one-out (LOO) cross-validation uses one data point in the original set as the assessment data and all other data points as the analysis set. A LOO resampling set has as many resamples as rows in the original data set.

Usage

```
loo_cv(data, ...)
```

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>data</td>
<td>A data frame.</td>
</tr>
<tr>
<td>...</td>
<td>Not currently used.</td>
</tr>
</tbody>
</table>

Value

An tibble with classes loo_cv, rset, tbl_df, tbl, and data.frame. The results include a column for the data split objects and one column called id that has a character string with the resample identifier.

Examples

```
loo_cv(mtcars)
```
Description

For stratified resampling, this function can create strata from numeric data and also make non-numeric data more conducive to be used for stratification.

Usage

make_strata(x, breaks = 4, nunique = 5, pool = 0.1, depth = 20)

Arguments

- **x**: An input vector.
- **breaks**: A single number giving the number of bins desired to stratify a numeric stratification variable.
- **nunique**: An integer for the number of unique value threshold in the algorithm.
- **pool**: A proportion of data used to determine if a particular group is too small and should be pooled into another group.
- **depth**: An integer that is used to determine the best number of percentiles that should be used. The number of bins are based on \( \min(5, \text{floor}(n / \text{depth})) \) where \( n = \text{length}(x) \). If \( x \) is numeric, there must be at least 40 rows in the data set (when \( \text{depth} = 20 \)) to conduct stratified sampling.

Details

For numeric data, if the number of unique levels is less than \( \text{nunique} \), the data are treated as categorical data.

For categorical inputs, the function will find levels of \( x \) that occur in the data with percentage less than \( \text{pool} \). The values from these groups will be randomly assigned to the remaining strata (as will data points that have missing values in \( x \)).

For numeric data with more unique values than \( \text{nunique} \), the data will be converted to being categorical based on percentiles of the data. The percentile groups will have no more than 20 percent of the data in each group. Again, missing values in \( x \) are randomly assigned to groups.

Value

A factor vector.
Examples

```r
set.seed(61)
x1 <- rpois(100, lambda = 5)
table(x1)
table(make_strata(x1))

set.seed(554)
x2 <- rpois(100, lambda = 1)
table(x2)
table(make_strata(x2))

# small groups are randomly assigned
x3 <- factor(x2)
table(x3)
table(make_strata(x3))

# `oilType` data from `caret`
x4 <- rep(LETTERS[1:7], c(37, 26, 3, 7, 11, 10, 2))
table(x4)
table(make_strata(x4))
table(make_strata(x4, pool = 0.1))
table(make_strata(x4, pool = 0.0))

# not enough data to stratify
x5 <- rnorm(20)
table(make_strata(x5))

set.seed(483)
x6 <- rnorm(200)
quantile(x6, probs = (0:10)/10)
table(make_strata(x6, breaks = 10))
```

---

**mc.cv**

*Monte Carlo Cross-Validation*

**Description**

One resample of Monte Carlo cross-validation takes a random sample (without replacement) of the original data set to be used for analysis. All other data points are added to the assessment set.

**Usage**

```r
mc_cv(data, prop = 3/4, times = 25, strata = NULL, breaks = 4, ...)
```

**Arguments**

- `data` A data frame.
- `prop` The proportion of data to be retained for modeling/analysis.
- `times` The number of times to repeat the sampling.
strata  A variable that is used to conduct stratified sampling to create the resamples. This could be a single character value or a variable name that corresponds to a variable that exists in the data frame.

breaks  A single number giving the number of bins desired to stratify a numeric stratification variable.

...  Not currently used.

Details

The `strata` argument causes the random sampling to be conducted within the stratification variable. This can help ensure that the number of data points in the analysis data is equivalent to the proportions in the original data set. (Strata below 10% of the total are pooled together.)

Value

An `tibble` with classes `mc_cv`, `rset`, `tbl_df`, `tbl`, and `data.frame`. The results include a column for the data split objects and a column called `id` that has a character string with the resample identifier.

Examples

```r
mc_cv(mtcars, times = 2)
microbenchmark(mc_cv(mtcars, prop = .5, times = 2))
library(purrr)
iris2 <- iris[1:130,]
set.seed(13)
resample1 <- mc_cv(iris2, times = 3, prop = .5)
map_dbl(resample1$splits,
function(x) {
  dat <- as.data.frame(x)$Species
  mean(dat == "virginica")
})
set.seed(13)
resample2 <- mc_cv(iris2, strata = "Species", times = 3, prop = .5)
map_dbl(resample2$splits,
function(x) {
  dat <- as.data.frame(x)$Species
  mean(dat == "virginica")
})
set.seed(13)
resample3 <- mc_cv(iris2, strata = "Sepal.Length", breaks = 6, times = 3, prop = .5)
map_dbl(resample3$splits,
function(x) {
  dat <- as.data.frame(x)$Species
  mean(dat == "virginica")
})
```
**nested_cv**

**Nested or Double Resampling**

**Description**

`nested_cv` can be used to take the results of one resampling procedure and conduct further resamples within each split. Any type of resampling used in `rsample` can be used.

**Usage**

`nested_cv(data, outside, inside)`

**Arguments**

- **data** A data frame.
- **outside** The initial resampling specification. This can be an already created object or an expression of a new object (see the examples below). If the latter is used, the `data` argument does not need to be specified and, if it is given, will be ignored.
- **inside** An expression for the type of resampling to be conducted within the initial procedure.

**Details**

It is a bad idea to use bootstrapping as the outer resampling procedure (see the example below)

**Value**

An tibble with classe `nested_cv` and any other classes that outer resampling process normally contains. The results include a column for the outer data split objects, one or more id columns, and a column of nested tibbles called `inner_resamples` with the additional resamples.

**Examples**

```r
## Using expressions for the resampling procedures:
nested_cv(mtcars, outside = vfold_cv(v = 3), inside = bootstraps(times = 5))

## Using an existing object:
folds <- vfold_cv(mtcars)
nested_cv(mtcars, folds, inside = bootstraps(times = 5))

## The dangers of outer bootstraps:
set.seed(2222)
bad_idea <- nested_cv(mtcars, outside = bootstraps(times = 5),
                       inside = vfold_cv(v = 3))

first_outer_split <- bad_idea$splits[[1]]
outer_analysis <- as.data.frame(first_outer_split)
```
sum(grepl("Volvo 142E", rownames(outer_analysis)))
## For the 3-fold CV used inside of each bootstrap, how are the replicated
## 'Volvo 142E' data partitioned?
first_inner_split <- bad_idea$inner_resamples[[1]]$splits[[1]]
inner_analysis <- as.data.frame(first_inner_split)
inner_assess <- as.data.frame(first_inner_split, data = "assessment")
sum(grepl("Volvo 142E", rownames(inner_analysis)))
sum(grepl("Volvo 142E", rownames(inner_assess)))

---

**populate**

*Add Assessment Indicies*

**Description**

Many `rsplit` and `rset` objects do not contain indicators for the assessment samples. `populate()` can be used to fill the slot for the appropriate indices.

**Usage**

`populate(x, ...)`

**Arguments**

- `x` A `rsplit` and `rset` object.
- `...` Not currently used

**Value**

An object of the same kind with the integer indicies.

**Examples**

```r
set.seed(28432)
fold_rs <- vfold_cv(mtcars)

fold_rs$splits[[1]]$out_id
complement(fold_rs$splits[[1]])

populate(fold_rs$splits[[1]])$out_id

fold_rs_all <- populate(fold_rs)
fold_rs_all$splits[[1]]$out_id
```
rolling_origin

Rolling Origin Forecast Resampling

Description

This resampling method is useful when the data set has a strong time component. The resamples are not random and contain data points that are consecutive values. The function assumes that the original data set are sorted in time order.

Usage

```r
rolling_origin(
  data,
  initial = 5,
  assess = 1,
  cumulative = TRUE,
  skip = 0,
  lag = 0,
  ...
)
```

Arguments

- `data` A data frame.
- `initial` The number of samples used for analysis/modeling in the initial resample.
- `assess` The number of samples used for each assessment resample.
- `cumulative` A logical. Should the analysis resample grow beyond the size specified by `initial` at each resample?
- `skip` A integer indicating how many (if any) additional resamples to skip to thin the total amount of data points in the analysis resample. See the example below.
- `lag` A value to include an lag between the assessment and analysis set. This is useful if lagged predictors will be used during training and testing.
- `...` Not currently used.

Details

The main options, `initial` and `assess`, control the number of data points from the original data that are in the analysis and assessment set, respectively. When `cumulative = TRUE`, the analysis set will grow as resampling continues while the assessment set size will always remain static. `skip` enables the function to not use every data point in the resamples. When `skip = 0`, the resampling data sets will increment by one position. Suppose that the rows of a data set are consecutive days. Using `skip = 6` will make the analysis data set operate on weeks instead of days. The assessment set size is not affected by this option.
Value

An tibble with classes rolling_origin, rset, tbl_df, tbl, and data.frame. The results include a column for the data split objects and a column called id that has a character string with the resample identifier.

Examples

```r
set.seed(1131)
ex_data <- data.frame(row = 1:20, some_var = rnorm(20))
dim(rolling_origin(ex_data))
dim(rolling_origin(ex_data, skip = 2))
dim(rolling_origin(ex_data, skip = 2, cumulative = FALSE))

# You can also roll over calendar periods by first nesting by that period, # which is especially useful for irregular series where a fixed window # is not useful. This example slides over 5 years at a time.
library(dplyr)
library(tidyr)
data(drinks, package = "modeldata")

drinks_annual <- drinks %>%
  mutate(year = as.POSIXlt(date)$year + 1900) %>%
  nest(-year)

multi_year_roll <- rolling_origin(drinks_annual, cumulative = FALSE)

analysis(multi_year_roll$splits[[1]])
assessment(multi_year_roll$splits[[1]])
```

rsample: General Resampling Infrastructure for R

Description

`rsample` has functions to create variations of a data set that can be used to evaluate models or to estimate the sampling distribution of some statistic.

Terminology

- A **resample** is the result of a two-way split of a data set. For example, when bootstrapping, one part of the resample is a sample with replacement of the original data. The other part of the split contains the instances that were not contained in the bootstrap sample. The data structure `rsplit` is used to store a single resample.

- When the data are split in two, the portion that are used to estimate the model or calculate the statistic is called the **analysis** set here. In machine learning this is sometimes called the "training set" but this would be poorly named since it might conflict with any initial split of the original data.
• Conversely, the other data in the split are called the **assessment** data. In bootstrapping, these data are often called the "out-of-bag" samples.

• A collection of resamples is contained in an **rset** object.

### Basic Functions

The main resampling functions are: `vfold_cv()`, `bootstraps()`, `mc_cv()`, `rolling_origin()`, and `nested_cv()`.

---

### Description

`rsample` should be fully compatible with `dplyr 1.0.0`. With older versions of `dplyr`, there is partial support for the following verbs: `mutate()`, `arrange()`, `filter()`, `rename()`, `select()`, and `slice()`. We strongly recommend updating to `dplyr 1.0.0` if possible to get more complete integration with `dplyr`.

### Version Specific Behavior

`rsample` performs somewhat differently depending on whether you have `dplyr >= 1.0.0` (new) or `dplyr < 1.0.0` (old). Additionally, version 0.0.7 of `rsample` (new) introduced some changes to how `rsample` objects work with `dplyr`, even on old `dplyr`. Most of these changes influence the return value of a `dplyr` verb and determine whether it will be a tibble or an `rsample rset` subclass.

The table below attempts to capture most of these changes. These examples are not exhaustive and may not capture some edge-cases.

<table>
<thead>
<tr>
<th align="center">Joins:</th>
<th align="center">The following affect all of the <code>dplyr</code> joins, such as <code>left_join()</code>, <code>right_join()</code>, <code>full_join()</code>, and <code>inner_join()</code>.</th>
<th align="center"><code>operation</code></th>
<th align="center"><code>old sample + old dplyr</code></th>
<th align="center"><code>new sample + old dplyr</code></th>
<th align="center"><code>new sample + new dplyr</code></th>
</tr>
</thead>
<tbody>
<tr>
<td align="center"><code>join(rset, tbl)</code></td>
<td align="center">error</td>
<td align="center">error</td>
<td align="center">tibble</td>
<td align="center"></td>
<td align="center"></td>
</tr>
</tbody>
</table>

The idea here is that, if there are less rows in the result, the result should not be an `rset` object. For example, you can’t have a 10-fold CV object without 10 rows.

Joins that keep the rows of the original `rset` object:

<table>
<thead>
<tr>
<th align="center">operation</th>
<th align="center"><code>old sample + old dplyr</code></th>
<th align="center"><code>new sample + old dplyr</code></th>
<th align="center"><code>new sample + new dplyr</code></th>
</tr>
</thead>
<tbody>
<tr>
<td align="center"><code>join(rset, tbl)</code></td>
<td align="center">error</td>
<td align="center">error</td>
<td align="center">rset</td>
</tr>
</tbody>
</table>

As with the logic above, if the original `rset` object (defined by the split column and the id col-
umn(s) is left intact, the results should be an rset.

Row Subsetting:
As mentioned above, this should result in a tibble if any rows are removed or added. Simply reordering rows still results in a valid rset with new rsample.
Cases where rows are removed or added:

<table>
<thead>
<tr>
<th>operation</th>
<th>old rsample + old dplyr</th>
<th>new rsample + old dplyr</th>
<th>new rsample + new dplyr</th>
</tr>
</thead>
<tbody>
<tr>
<td>rset[ind,]</td>
<td>tibble</td>
<td>tibble</td>
<td>tibble</td>
</tr>
<tr>
<td>slice(rset)</td>
<td>rset</td>
<td>tibble</td>
<td>tibble</td>
</tr>
<tr>
<td>filter(rset)</td>
<td>rset</td>
<td>tibble</td>
<td>tibble</td>
</tr>
</tbody>
</table>

Cases where all rows are kept, but are possibly reordered:

<table>
<thead>
<tr>
<th>operation</th>
<th>old rsample + old dplyr</th>
<th>new rsample + old dplyr</th>
<th>new rsample + new dplyr</th>
</tr>
</thead>
<tbody>
<tr>
<td>rset[ind,]</td>
<td>tibble</td>
<td>rset</td>
<td>rset</td>
</tr>
<tr>
<td>slice(rset)</td>
<td>rset</td>
<td>rset</td>
<td>rset</td>
</tr>
<tr>
<td>filter(rset)</td>
<td>rset</td>
<td>rset</td>
<td>rset</td>
</tr>
<tr>
<td>arrange(rset)</td>
<td>rset</td>
<td>rset</td>
<td>rset</td>
</tr>
</tbody>
</table>

Column Subsetting:

When the splits column or any id columns are dropped or renamed, the result should no longer be considered a valid rset.
Cases when the required columns are removed or renamed:

<table>
<thead>
<tr>
<th>operation</th>
<th>old rsample + old dplyr</th>
<th>new rsample + old dplyr</th>
<th>new rsample + new dplyr</th>
</tr>
</thead>
<tbody>
<tr>
<td>rset[,ind]</td>
<td>tibble</td>
<td>tibble</td>
<td>tibble</td>
</tr>
<tr>
<td>select(rset)</td>
<td>rset</td>
<td>tibble</td>
<td>tibble</td>
</tr>
<tr>
<td>rename(rset)</td>
<td>tibble</td>
<td>tibble</td>
<td>tibble</td>
</tr>
</tbody>
</table>

Cases when no required columns are affected:

<table>
<thead>
<tr>
<th>operation</th>
<th>old rsample + old dplyr</th>
<th>new rsample + old dplyr</th>
<th>new rsample + new dplyr</th>
</tr>
</thead>
<tbody>
<tr>
<td>rset[,ind]</td>
<td>tibble</td>
<td>rset</td>
<td>rset</td>
</tr>
<tr>
<td>select(rset)</td>
<td>rset</td>
<td>rset</td>
<td>rset</td>
</tr>
<tr>
<td>rename(rset)</td>
<td>rset</td>
<td>rset</td>
<td>rset</td>
</tr>
</tbody>
</table>

Other Column Operations:

Cases when the required columns are altered:

<table>
<thead>
<tr>
<th>operation</th>
<th>old rsample + old dplyr</th>
<th>new rsample + old dplyr</th>
<th>new rsample + new dplyr</th>
</tr>
</thead>
<tbody>
<tr>
<td>mutate(rset)</td>
<td>rset</td>
<td>tibble</td>
<td>tibble</td>
</tr>
</tbody>
</table>
Cases when no required columns are affected:

<table>
<thead>
<tr>
<th>operation</th>
<th>old rsample + old dplyr</th>
<th>new rsample + old dplyr</th>
<th>new rsample + new dplyr</th>
</tr>
</thead>
<tbody>
<tr>
<td>mutate(rset)</td>
<td>rset</td>
<td>rset</td>
<td>rset</td>
</tr>
</tbody>
</table>

rsample2caret  

Convert Resampling Objects to Other Formats

Description

These functions can convert resampling objects between `rsample` and `caret`.

Usage

```r
rsample2caret(object, data = c("analysis", "assessment"))
caret2rsample(ctrl, data = NULL)
```

Arguments

- `object`: An rset object. Currently, nested_cv is not supported.
- `data`: The data that was originally used to produce the ctrl object.
- `ctrl`: An object produced by trainControl that has had the index and indexOut elements populated by integers. One method of getting this is to extract the control objects from an object produced by `train`.

Value

- `rsample2caret` returns a list that mimics the index and indexOut elements of a `trainControl` object. `caret2rsample` returns an rset object of the appropriate class.

tidy.rsplit  

Tidy Resampling Object

Description

The tidy function from the `broom` package can be used on rset and rsplit objects to generate tibbles with which rows are in the analysis and assessment sets.
Usage

## S3 method for class 'rsplit'
tidy(x, unique_ind = TRUE, ...)

## S3 method for class 'rset'
tidy(x, ...)

## S3 method for class 'vfold_cv'
tidy(x, ...)

## S3 method for class 'nested_cv'
tidy(x, ...)

Arguments

x A rset or rsplit object

unique_ind Should unique row identifiers be returned? For example, if FALSE then bootstrapping results will include multiple rows in the sample for the same row in the original data.

... Not currently used.

Details

Note that for nested resampling, the rows of the inner resample, named inner_Row, are relative row indices and do not correspond to the rows in the original data set.

Value

A tibble with columns Row and Data. The latter has possible values "Analysis" or "Assessment".

For rset inputs, identification columns are also returned but their names and values depend on the type of resampling. vfold_cv contains a column "Fold" and, if repeats are used, another called "Repeats". bootstraps and mc_cv use the column "Resample".

Examples

library(ggplot2)
theme_set(theme_bw())

set.seed(4121)
cv <- tidy(vfold_cv(mtcars, v = 5))
ggplot(cv, aes(x = Fold, y = Row, fill = Data)) +
  geom_tile() + scale_fill_brewer()

set.seed(4121)
rcv <- tidy(vfold_cv(mtcars, v = 5, repeats = 2))
ggplot(rcv, aes(x = Fold, y = Row, fill = Data)) +
  geom_tile() + facet_wrap(~Repeat) + scale_fill_brewer()

set.seed(4121)
Two Class Data

Description

Two Class Data

Details

There are artificial data with two predictors (A and B) and a factor outcome variable (Class). These data are now in the modeldata package.

validation_split

Create a Validation Set

Description

validation_split() takes a single random sample (without replacement) of the original data set to be used for analysis. All other data points are added to the assessment set (to be used as the validation set).

Usage

validation_split(data, prop = 3/4, strata = NULL, breaks = 4, ...)

mccv <- tidy(mc_cv(mtcars, times = 5))
ggplot(mccv, aes(x = Resample, y = Row, fill = Data)) + geom_tile() + scale_fill_brewer()

set.seed(4121)
btt <- tidy(bootstraps(mtcars, time = 5))
ggplot(bt, aes(x = Resample, y = Row, fill = Data)) + geom_tile() + scale_fill_brewer()

dat <- data.frame(day = 1:30)
# Resample by week instead of day
sts_cv <- rolling_origin(dat, initial = 7, assess = 7,
                          skip = 6, cumulative = FALSE)
sts_cv <- tidy(sts_cv)
ggplot(sts_cv, aes(x = Resample, y = factor(Row), fill = Data)) + geom_tile() + scale_fill_brewer()
Arguments

- **data**: A data frame.
- **prop**: The proportion of data to be retained for modeling/analysis.
- **strata**: A variable that is used to conduct stratified sampling to create the resamples. This could be a single character value or a variable name that corresponds to a variable that exists in the data frame.
- **breaks**: A single number giving the number of bins desired to stratify a numeric stratification variable.
- **...**: Not currently used.

Details

The `strata` argument causes the random sampling to be conducted *within the stratification variable*. This can help ensure that the number of data points in the analysis data is equivalent to the proportions in the original data set. (Strata below 10% of the total are pooled together.)

Value

An tibble with classes `validation_split`, `rset`, `tbl_df`, `tbl`, and `data.frame`. The results include a column for the data split objects and a column called `id` that has a character string with the resample identifier.

Examples

```r
validation_split(iris, prop = .9)
```

---

**vfold_cv**  

**V-Fold Cross-Validation**

Description

V-fold cross-validation randomly splits the data into V groups of roughly equal size (called "folds"). A resample of the analysis data consisted of V-1 of the folds while the assessment set contains the final fold. In basic V-fold cross-validation (i.e. no repeats), the number of resamples is equal to V.

Usage

```r
vfold_cv(data, v = 10, repeats = 1, strata = NULL, breaks = 4, ...)
```

Arguments

- **data**: A data frame.
- **v**: The number of partitions of the data set.
- **repeats**: The number of times to repeat the V-fold partitioning.
strata

A variable that is used to conduct stratified sampling to create the folds. This
could be a single character value or a variable name that corresponds to a vari-
able that exists in the data frame.

breaks

A single number giving the number of bins desired to stratify a numeric stratifi-
cation variable.

... 

Not currently used.

Details

The strata argument causes the random sampling to be conducted within the stratification vari-
able. This can help ensure that the number of data points in the analysis data is equivalent to the
proportions in the original data set. (Strata below 10% of the total are pooled together.) When more
than one repeat is requested, the basic V-fold cross-validation is conducted each time. For example,
if three repeats are used with \( v = 10 \), there are a total of 30 splits which as three groups of 10 that
are generated separately.

Value

A tibble with classes vfold_cv, rset, tbl_df, tbl, and data.frame. The results include a column
for the data split objects and one or more identification variables. For a single repeats, there will be
one column called id that has a character string with the fold identifier. For repeats, id is the repeat
number and an additional column called id2 that contains the fold information (within repeat).

Examples

vfold_cv(mtcars, v = 10)
vfold_cv(mtcars, v = 10, repeats = 2)

library(purrr)
iris2 <- iris[1:130, ]

set.seed(13)
folds1 <- vfold_cv(iris2, v = 5)
map_dbl(folds1$splits,
    function(x) {
        dat <- as.data.frame(x)$Species
        mean(dat == "virginica")
    })

set.seed(13)
folds2 <- vfold_cv(iris2, strata = "Species", v = 5)
map_dbl(folds2$splits,
    function(x) {
        dat <- as.data.frame(x)$Species
        mean(dat == "virginica")
    })

set.seed(13)
folds3 <- vfold_cv(iris2, strata = "Petal.Length", breaks = 6, v = 5)
map_dbl(folds3$splits,
    function(x) {
dat <- as.data.frame(x)$Species
mean(dat == "virginica")
}

vfold_cv
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