# Package 'cvms'

January 14, 2025

Title Cross-Validation for Model Selection

Version 1.6.3

Description Cross-validate one or multiple regression and classification models and get relevant evaluation metrics in a tidy format. Validate the best model on a test set and compare it to a baseline evaluation.
Alternatively, evaluate predictions from an external model. Currently supports regression and classification (binary and multiclass).
Described in chp. 5 of Jeyaraman, B. P., Olsen, L. R.,
& Wambugu M. (2019, ISBN: 9781838550134).

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URL https://github.com/ludvigolsen/cvms

BugReports https://github.com/ludvigolsen/cvms/issues

**Depends** R (>= 3.5)

Imports checkmate (>= 2.0.0), data.table (>= 1.12), dplyr (>= 0.8.5), ggplot2, groupdata2 (>= 2.0.5), lifecycle, lme4 (>= 1.1-23), MuMIn (>= 1.43.17), parameters (>= 0.15.0), plyr, pROC (>= 1.16.0), purrr, rearrr (>= 0.3.4), recipes (>= 0.1.13), rlang (>= 0.4.7), stats, stringr, tibble (>= 3.0.3), tidyr (>= 1.1.2), utils

**Suggests** AUC, covr (>= 3.3.1), e1071 (>= 1.7-2), furrr, ggimage (>= 0.3.3), ggnewscale (>= 0.5.0), knitr, merDeriv (>= 0.2-4), nnet (>= 7.3-20), randomForest (>= 4.6-14), rmarkdown, rsvg, testthat (>= 2.3.2), xpectr (>= 0.4.1)

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## **Description**

cvms-package

Perform (repeated) cross-validation on a list of model formulas. Validate the best model on a validation set. Perform baseline evaluations on your test set. Generate model formulas by combining your fixed effects. Evaluate predictions from an external model.

#### **Details**

Returns results in a tibble for easy comparison, reporting and further analysis.

The main functions are: cross\_validate(), cross\_validate\_fn(), validate(), validate\_fn(), baseline(), and evaluate().

#### Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

## See Also

Useful links:

- https://github.com/ludvigolsen/cvms
- Report bugs at https://github.com/ludvigolsen/cvms/issues

baseline Create baseline evaluations

## Description

#### [Maturing]

Create a baseline evaluation of a test set.

In modelling, a *baseline* is a result that is meaningful to compare the results from our models to. For instance, in classification, we usually want our results to be better than *random guessing*. E.g. if we have three classes, we can expect an accuracy of 33.33%, as for every observation we have 1/3 chance of guessing the correct class. So our model should achieve a higher accuracy than 33.33% before it is more useful to us than guessing.

While this expected value is often fairly straightforward to find analytically, it only represents what we can expect on average. In reality, it's possible to get far better results than that by guessing. baseline() (binomial, multinomial) finds the range of likely values by evaluating multiple sets

of random predictions and summarizing them with a set of useful descriptors. If random guessing frequently obtains an accuracy of 40%, perhaps our model should have better performance than this, before we declare it better than guessing.

#### How:

When `family` is binomial: evaluates `n` sets of random predictions against the dependent variable, along with a set of all 0 predictions and a set of all 1 predictions. See also baseline\_binomial().

When `family` is multinomial: creates *one-vs-all* (binomial) baseline evaluations for `n` sets of random predictions against the dependent variable, along with sets of "all class x,y,z,..." predictions. See also baseline\_multinomial().

When `family` is gaussian: fits baseline models  $(y \sim 1)$  on `n` random subsets of `train\_data` and evaluates each model on `test\_data`. Also evaluates a model fitted on all rows in `train\_data`. See also baseline\_gaussian().

## Wrapper functions:

Consider using one of the wrappers, as they are simpler to use and understand: baseline\_gaussian(), baseline\_multinomial(), and baseline\_binomial().

#### Usage

```
baseline(
  test_data,
  dependent_col,
  family,
  train_data = NULL,
  n = 100,
 metrics = list(),
 positive = 2,
  cutoff = 0.5,
  random_generator_fn = runif,
  random_effects = NULL,
 min_training_rows = 5,
 min_training_rows_left_out = 3,
 REML = FALSE,
 parallel = FALSE
)
```

#### **Arguments**

dependent\_col Name of dependent variable in the supplied test and training sets.

Name of family. (Character)
Currently supports "gaussian", "binomial" and "multinomial".

train\_data data.frame. Only used when `family` is "gaussian".

Number of random samplings to perform. (Default is 100)
For gaussian: The number of random samplings of `train\_data` to fit baseline models on.

For binomial and multinomial: The number of sets of random predictions to evaluate.

metrics

list for enabling/disabling metrics.

E.g. list("RMSE" = FALSE) would remove RMSE from the regression results, and list("Accuracy" = TRUE) would add the regular Accuracy metric to the classification results. Default values (TRUE/FALSE) will be used for the remaining available metrics.

You can enable/disable all metrics at once by including "all" = TRUE/FALSE in the list. This is done prior to enabling/disabling individual metrics, why f.i. list("all" = FALSE, "RMSE" = TRUE) would return only the RMSE metric.

The list can be created with gaussian\_metrics(), binomial\_metrics(), or multinomial\_metrics().

Also accepts the string "all".

positive

Level from dependent variable to predict. Either as character (*preferable*) or level index (1 or 2 - alphabetically).

E.g. if we have the levels "cat" and "dog" and we want "dog" to be the positive class, we can either provide "dog" or 2, as alphabetically, "dog" comes after "cat".

**Note:** For *reproducibility*, it's preferable to **specify the name directly**, as different locales may sort the levels differently.

Used when calculating confusion matrix metrics and creating ROC curves.

N.B. Only affects evaluation metrics, not the returned predictions.

N.B. **Binomial only**. (Character or Integer)

cutoff

Threshold for predicted classes. (Numeric)

N.B. Binomial only

random\_generator\_fn

Function for generating random numbers when type is "multinomial". The softmax function is applied to the generated numbers to transform them to probabilities.

The first argument must be the number of random numbers to generate, as no other arguments are supplied.

To test the effect of using different functions, see multiclass\_probability\_tibble().

N.B. Multinomial only

random\_effects Random effects structure for the Gaussian baseline model. (Character)

E.g. with "(1|ID)", the model becomes " $y \sim 1 + (1|ID)$ ".

N.B. Gaussian only

min\_training\_rows

Minimum number of rows in the random subsets of `train\_data`.

Gaussian only. (Integer)

min\_training\_rows\_left\_out

Minimum number of rows left out of the random subsets of `train\_data`.

I.e. a subset will maximally have the size:

max\_rows\_in\_subset = nrow(`train\_data`) - `min\_training\_rows\_left\_out`.

N.B. Gaussian only. (Integer)

REML Whether to use Restricted Maximum Likelihood. (Logical)

N.B. Gaussian only. (Integer)

parallel Whether to run the `n` evaluations in parallel. (Logical)

Remember to register a parallel backend first. E.g. with doParallel::registerDoParallel.

#### **Details**

## Packages used:

## **Models:**

Gaussian: stats::lm, lme4::lmer

#### **Results:** Gaussian:

r2m: MuMIn::r.squaredGLMMr2c: MuMIn::r.squaredGLMM

AIC:stats::AIC
AICc:MuMIn::AICc
BIC:stats::BIC

## Binomial and Multinomial:

ROC and related metrics: Binomial: pROC::roc

Multinomial: pROC::multiclass.roc

## Value

list containing:

- 1. a tibble with summarized results (called summarized\_metrics)
- 2. a tibble with random evaluations (random\_evaluations)
- 3. a tibble with the summarized class level results (summarized\_class\_level\_results) (Multinomial only)

## **Gaussian Results:**

The **Summarized Results** tibble contains:

Average RMSE, MAE, NRMSE(IQR), RRSE, RAE, RMSLE.

See the additional metrics (disabled by default) at ?gaussian\_metrics.

The **Measure** column indicates the statistical descriptor used on the evaluations. The row where Measure == All\_rows is the evaluation when the baseline model is trained on all rows in `train\_data`.

The **Training Rows** column contains the aggregated number of rows used from `train\_data`, when fitting the baseline models.

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The Random Evaluations tibble contains:

The non-aggregated metrics.

A nested tibble with the **predictions** and targets.

A nested tibble with the **coefficients** of the baseline models.

Number of training rows used when fitting the baseline model on the training set.

A nested **Process** information object with information about the evaluation.

Name of **dependent** variable.

Name of **fixed** effect (bias term only).

Random effects structure (if specified).

**Binomial Results:** 

Based on the generated test set predictions, a confusion matrix and ROC curve are used to get the following:

ROC:

AUC, Lower CI, and Upper CI

Note, that the ROC curve is only computed when AUC is enabled.

Confusion Matrix:

Balanced Accuracy, Accuracy, F1, Sensitivity, Specificity, Positive Predictive Value, Negative Predictive Value, Kappa, Detection Rate, Detection Prevalence, Prevalence, and MCC (Matthews correlation coefficient).

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#### The **Summarized Results** tibble contains:

The **Measure** column indicates the statistical descriptor used on the evaluations. The row where Measure == All\_0 is the evaluation when all predictions are 0. The row where Measure == All\_1 is the evaluation when all predictions are 1.

The aggregated metrics.

.....

The Random Evaluations tibble contains:

The non-aggregated metrics.

A nested tibble with the **predictions** and targets.

A list of **ROC** curve objects (if computed).

A nested tibble with the **confusion matrix**. The Pos\_ columns tells you whether a row is a True Positive (TP), True Negative (TN), False Positive (FP), or False Negative (FN), depending on which level is the "positive" class. I.e. the level you wish to predict.

A nested **Process** information object with information about the evaluation.

Name of **dependent** variable.

#### **Multinomial Results:**

Based on the generated test set predictions, one-vs-all (binomial) evaluations are performed and aggregated to get the same metrics as in the binomial results (excluding MCC, AUC, Lower CI and Upper CI), with the addition of **Overall Accuracy** and *multiclass* **MCC** in the summarized

results. It is possible to enable multiclass **AUC** as well, which has been disabled by default as it is slow to calculate when there's a large set of classes.

Since we use macro-averaging, Balanced Accuracy is the macro-averaged metric, *not* the macro sensitivity as sometimes used.

Note: we also refer to the *one-vs-all evaluations* as the *class level results*.

.....

#### The Summarized Results tibble contains:

Summary of the random evaluations.

**How**: First, the one-vs-all binomial evaluations are aggregated by repetition, then, these aggregations are summarized. Besides the metrics from the binomial evaluations (see *Binomial Results* above), it also includes Overall Accuracy and *multiclass* MCC.

The Measure column indicates the statistical descriptor used on the evaluations. The Mean, Median, SD, IQR, Max, Min, NAs, and INFs measures describe the *Random Evaluations* tibble, while the CL Max, CL Min, CL NAs, and CL INFs describe the Class Level results.

The rows where Measure == All\_<<class name>> are the evaluations when all the observations are predicted to be in that class.

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#### The **Summarized Class Level Results** tibble contains:

The (nested) summarized results for each class, with the same metrics and descriptors as the *Summarized Results* tibble. Use tidyr::unnest on the tibble to inspect the results.

**How**: The one-vs-all evaluations are summarized by class.

The rows where Measure == All\_0 are the evaluations when none of the observations are predicted to be in that class, while the rows where Measure == All\_1 are the evaluations when all of the observations are predicted to be in that class.

.....

#### The Random Evaluations tibble contains:

The repetition results with the same metrics as the *Summarized Results* tibble.

**How**: The one-vs-all evaluations are aggregated by repetition. If a metric contains one or more NAs in the one-vs-all evaluations, it will lead to an NA result for that repetition.

Also includes:

A nested tibble with the one-vs-all binomial evaluations (**Class Level Results**), including nested **Confusion Matrices** and the **Support** column, which is a count of how many observations from the class is in the test set.

A nested tibble with the **predictions** and targets.

A list of **ROC** curve objects.

A nested tibble with the multiclass confusion matrix.

A nested **Process** information object with information about the evaluation.

Name of **dependent** variable.

#### Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

## See Also

Other baseline functions: baseline\_binomial(), baseline\_gaussian(), baseline\_multinomial()

## **Examples**

```
# Attach packages
library(cvms)
library(groupdata2) # partition()
library(dplyr) # %>% arrange()
library(tibble)
# Data is part of cvms
data <- participant.scores</pre>
# Set seed for reproducibility
set.seed(1)
# Partition data
partitions <- partition(data, p = 0.7, list_out = TRUE)</pre>
train_set <- partitions[[1]]</pre>
test_set <- partitions[[2]]</pre>
# Create baseline evaluations
# Note: usually n=100 is a good setting
# Gaussian
baseline(
  test_data = test_set, train_data = train_set,
  dependent_col = "score", random_effects = "(1|session)",
  n = 2, family = "gaussian"
)
# Binomial
baseline(
  test_data = test_set, dependent_col = "diagnosis",
  n = 2, family = "binomial"
# Multinomial
# Create some data with multiple classes
multiclass_data <- tibble(</pre>
  "target" = rep(paste0("class_", 1:5), each = 10)
) %>%
  dplyr::sample_n(35)
baseline(
  test_data = multiclass_data,
  dependent_col = "target",
  n = 4, family = "multinomial"
# Parallelize evaluations
# Attach doParallel and register four cores
# Uncomment:
```

```
# library(doParallel)
# registerDoParallel(4)
# Binomial
baseline(
  test_data = test_set, dependent_col = "diagnosis",
  n = 4, family = "binomial"
  #, parallel = TRUE # Uncomment
)
# Gaussian
baseline(
  test_data = test_set, train_data = train_set,
  dependent_col = "score", random_effects = "(1|session)",
  n = 4, family = "gaussian"
  #, parallel = TRUE # Uncomment
)
# Multinomial
(mb <- baseline(</pre>
  test_data = multiclass_data,
  dependent_col = "target",
  n = 6, family = "multinomial"
  #, parallel = TRUE # Uncomment
))
# Inspect the summarized class level results
# for class_2
mb$summarized_class_level_results %>%
  dplyr::filter(Class == "class_2") %>%
  tidyr::unnest(Results)
# Multinomial with custom random generator function
# that creates very "certain" predictions
# (once softmax is applied)
rcertain <- function(n) {</pre>
  (runif(n, min = 1, max = 100)^1.4) / 100
baseline(
  test_data = multiclass_data,
  dependent_col = "target",
  n = 6, family = "multinomial",
  random_generator_fn = rcertain
  #, parallel = TRUE # Uncomment
)
```

#### **Description**

## [Maturing]

Create a baseline evaluation of a test set.

In modelling, a *baseline* is a result that is meaningful to compare the results from our models to. For instance, in classification, we usually want our results to be better than *random guessing*. E.g. if we have three classes, we can expect an accuracy of 33.33%, as for every observation we have 1/3 chance of guessing the correct class. So our model should achieve a higher accuracy than 33.33% before it is more useful to us than guessing.

While this expected value is often fairly straightforward to find analytically, it only represents what we can expect on average. In reality, it's possible to get far better results than that by guessing. baseline\_binomial() finds the range of likely values by evaluating multiple sets of random predictions and summarizing them with a set of useful descriptors. Additionally, it evaluates a set of all 0 predictions and a set of all 1 predictions.

## Usage

```
baseline_binomial(
  test_data,
  dependent_col,
  n = 100,
  metrics = list(),
  positive = 2,
  cutoff = 0.5,
  parallel = FALSE
)
```

#### **Arguments**

test\_data data.frame.

dependent\_col Name of dependent variable in the supplied test and training sets.

n The number of sets of random predictions to evaluate. (Default is 100)

metrics list for enabling/disabling metrics.

E.g. list("F1" = FALSE) would remove F1 from the results, and list("Accuracy" = TRUE) would add the regular Accuracy metric to the results. Default values (TRUE/FALSE) will be used for the remaining available metrics.

You can enable/disable all metrics at once by including "all" = TRUE/FALSE in the list. This is done prior to enabling/disabling individual metrics, why f.i. list("all" = FALSE, "Accuracy" = TRUE) would return only the Accuracy metric.

The list can be created with binomial\_metrics().

Also accepts the string "all".

positive Level from dependent variable to predict. Either as character (preferable) or

level index (1 or 2 - alphabetically).

E.g. if we have the levels "cat" and "dog" and we want "dog" to be the positive class, we can either provide "dog" or 2, as alphabetically, "dog" comes after "cat".

Note: For reproducibility, it's preferable to specify the name directly, as dif-

ferent locales may sort the levels differently.

Used when calculating confusion matrix metrics and creating ROC curves.

N.B. Only affects evaluation metrics, not the returned predictions.

cutoff Threshold for predicted classes. (Numeric)

parallel Whether to run the `n` evaluations in parallel. (Logical)

Remember to register a parallel backend first. E.g. with doParallel::registerDoParallel.

#### **Details**

Packages used:

ROC and AUC: pROC::roc

#### Value

list containing:

1. a tibble with summarized results (called summarized\_metrics)

2. a tibble with random evaluations (random\_evaluations)

Decid on the control test and disting a confession on

Based on the generated test set predictions, a confusion matrix and ROC curve are used to get the following:

ROC:

AUC, Lower CI, and Upper CI

Note, that the ROC curve is only computed when AUC is enabled.

Confusion Matrix:

Balanced Accuracy, Accuracy, F1, Sensitivity, Specificity, Positive Predictive Value, Negative Predictive Value, Kappa, Detection Rate, Detection Prevalence, Prevalence, and MCC (Matthews correlation coefficient).

.....

## The Summarized Results tibble contains:

The **Measure** column indicates the statistical descriptor used on the evaluations. The row where Measure == All\_0 is the evaluation when all predictions are 0. The row where Measure == All\_1 is the evaluation when all predictions are 1.

The aggregated metrics.

.....

The **Random Evaluations** tibble contains:

The non-aggregated metrics.

A nested tibble with the **predictions** and targets.

A list of **ROC** curve objects (if computed).

A nested tibble with the **confusion matrix**. The Pos\_ columns tells you whether a row is a True Positive (TP), True Negative (TN), False Positive (FP), or False Negative (FN), depending on which level is the "positive" class. I.e. the level you wish to predict.

A nested **Process** information object with information about the evaluation.

Name of **dependent** variable.

## Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

#### See Also

Other baseline functions: baseline(), baseline\_gaussian(), baseline\_multinomial()

## **Examples**

```
# Attach packages
library(cvms)
library(groupdata2) # partition()
library(dplyr) # %>% arrange()
# Data is part of cvms
data <- participant.scores</pre>
# Set seed for reproducibility
set.seed(1)
# Partition data
partitions <- partition(data, p = 0.7, list_out = TRUE)</pre>
train_set <- partitions[[1]]</pre>
test_set <- partitions[[2]]</pre>
# Create baseline evaluations
# Note: usually n=100 is a good setting
baseline_binomial(
  test_data = test_set,
  dependent_col = "diagnosis",
  n = 2
)
# Parallelize evaluations
# Attach doParallel and register four cores
# Uncomment:
# library(doParallel)
# registerDoParallel(4)
# Make sure to uncomment the parallel argument
baseline_binomial(
  test_data = test_set,
```

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```
dependent_col = "diagnosis",
  n = 4
  #, parallel = TRUE  # Uncomment
)
```

baseline\_gaussian

Create baseline evaluations for regression models

## Description

## [Maturing]

Create a baseline evaluation of a test set.

In modelling, a *baseline* is a result that is meaningful to compare the results from our models to. In regression, we want our model to be better than a model without any predictors. If our model does not perform better than such a simple model, it's unlikely to be useful.

baseline\_gaussian() fits the intercept-only model ( $y \sim 1$ ) on `n` random subsets of `train\_data` and evaluates each model on `test\_data`. Additionally, it evaluates a model fitted on all rows in `train\_data`.

## Usage

```
baseline_gaussian(
  test_data,
  train_data,
  dependent_col,
  n = 100,
  metrics = list(),
  random_effects = NULL,
  min_training_rows = 5,
  min_training_rows_left_out = 3,
  REML = FALSE,
  parallel = FALSE
)
```

## **Arguments**

```
train_data data.frame.

train_data data.frame.

dependent_col Name of dependent variable in the supplied test and training sets.

The number of random samplings of `train_data` to fit baseline models on.

(Default is 100)
```

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metrics list for enabling/disabling metrics.

E.g. list("RMSE" = FALSE) would remove RMSE from the results, and list("TAE" = TRUE) would add the Total Absolute Error metric to the results. Default

values (TRUE/FALSE) will be used for the remaining available metrics.

You can enable/disable all metrics at once by including "all" = TRUE/FALSE in the list. This is done prior to enabling/disabling individual metrics, why f.i. list("all" = FALSE, "RMSE" = TRUE) would return only the RMSE metric.

The list can be created with gaussian\_metrics().

Also accepts the string "all".

random\_effects Random effects structure for the baseline model. (Character)

E.g. with "(1|ID)", the model becomes " $y \sim 1 + (1|ID)$ ".

min\_training\_rows

Minimum number of rows in the random subsets of `train\_data`.

min\_training\_rows\_left\_out

Minimum number of rows left out of the random subsets of `train\_data`.

I.e. a subset will maximally have the size:

max\_rows\_in\_subset = nrow(`train\_data`) - `min\_training\_rows\_left\_out`.

REML Whether to use Restricted Maximum Likelihood. (Logical) parallel Whether to run the `n` evaluations in parallel. (Logical)

Remember to register a parallel backend first. E.g. with doParallel::registerDoParallel.

#### **Details**

Packages used:

## **Models:**

stats::lm, lme4::lmer

#### **Results:**

r2m: MuMIn::r.squaredGLMM r2c:MuMIn::r.squaredGLMM

AIC: stats::AIC
AICc: MuMIn::AICc
BIC: stats::BIC

## Value

list containing:

- 1. a tibble with summarized results (called summarized\_metrics)
- 2. a tibble with random evaluations (random\_evaluations)

•••••

The Summarized Results tibble contains:

Average RMSE, MAE, NRMSE(IQR), RRSE, RAE, RMSLE.

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See the additional metrics (disabled by default) at ?gaussian\_metrics.

The **Measure** column indicates the statistical descriptor used on the evaluations. The row where Measure == All\_rows is the evaluation when the baseline model is trained on all rows in `train\_data`.

The **Training Rows** column contains the aggregated number of rows used from `train\_data`, when fitting the baseline models.

.....

The **Random Evaluations** tibble contains:

The non-aggregated metrics.

A nested tibble with the **predictions** and targets.

A nested tibble with the **coefficients** of the baseline models.

Number of training rows used when fitting the baseline model on the training set.

A nested **Process** information object with information about the evaluation.

Name of **dependent** variable.

Name of **fixed** effect (bias term only).

Random effects structure (if specified).

#### Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

#### See Also

Other baseline functions: baseline(), baseline\_binomial(), baseline\_multinomial()

#### **Examples**

```
# Attach packages
library(cvms)
library(groupdata2) # partition()
library(dplyr) # %>% arrange()
# Data is part of cvms
data <- participant.scores
# Set seed for reproducibility
set.seed(1)
# Partition data
partitions <- partition(data, p = 0.7, list_out = TRUE)</pre>
train_set <- partitions[[1]]</pre>
test_set <- partitions[[2]]
# Create baseline evaluations
# Note: usually n=100 is a good setting
baseline_gaussian(
  test_data = test_set,
```

```
train_data = train_set,
 dependent_col = "score";
 random_effects = "(1|session)",
 n = 2
)
# Parallelize evaluations
# Attach doParallel and register four cores
# Uncomment:
# library(doParallel)
# registerDoParallel(4)
# Make sure to uncomment the parallel argument
baseline_gaussian(
 test_data = test_set,
 train_data = train_set,
 dependent_col = "score",
 random_effects = "(1|session)",
 n = 4
 #, parallel = TRUE # Uncomment
)
```

baseline\_multinomial Create baseline evaluations

#### **Description**

#### [Maturing]

Create a baseline evaluation of a test set.

In modelling, a *baseline* is a result that is meaningful to compare the results from our models to. For instance, in classification, we usually want our results to be better than *random guessing*. E.g. if we have three classes, we can expect an accuracy of 33.33%, as for every observation we have 1/3 chance of guessing the correct class. So our model should achieve a higher accuracy than 33.33% before it is more useful to us than guessing.

While this expected value is often fairly straightforward to find analytically, it only represents what we can expect on average. In reality, it's possible to get far better results than that by guessing. baseline\_multinomial() finds the range of likely values by evaluating multiple sets of random predictions and summarizing them with a set of useful descriptors.

Technically, it creates *one-vs-all* (binomial) baseline evaluations for the `n` sets of random predictions and summarizes them. Additionally, sets of "all class x,y,z,..." predictions are evaluated.

#### Usage

```
baseline_multinomial(
  test_data,
  dependent_col,
```

```
n = 100,
metrics = list(),
random_generator_fn = runif,
parallel = FALSE
)
```

#### **Arguments**

test\_data data.frame.

dependent\_col Name of dependent variable in the supplied test and training sets.

n The number of sets of random predictions to evaluate. (Default is 100)

metrics list for enabling/disabling metrics.

E.g. list("F1" = FALSE) would remove F1 from the results, and list("Accuracy" = TRUE) would add the regular Accuracy metric to the results. Default values

(TRUE/FALSE) will be used for the remaining available metrics.

You can enable/disable all metrics at once by including "all" = TRUE/FALSE in the list. This is done prior to enabling/disabling individual metrics, why f.i. list("all" = FALSE, "Accuracy" = TRUE) would return only the Accuracy

metric.

The list can be created with multinomial\_metrics().

Also accepts the string "all".

random\_generator\_fn

Function for generating random numbers. The softmax function is applied to the generated numbers to transform them to probabilities.

The first argument must be the number of random numbers to generate, as no

other arguments are supplied.

parallel Whether to run the `n` evaluations in parallel. (Logical)

Remember to register a parallel backend first. E.g. with doParallel::registerDoParallel.

To test the effect of using different functions, see multiclass\_probability\_tibble().

## Details

Packages used:

Multiclass ROC curve and AUC: pROC::multiclass.roc

## Value

list containing:

- 1. a tibble with summarized results (called summarized\_metrics)
- 2. a tibble with random evaluations (random\_evaluations)
- 3. a tibble with the summarized class level results (summarized\_class\_level\_results)

.....

#### **Macro metrics:**

Based on the generated predictions, *one-vs-all* (binomial) evaluations are performed and aggregated to get the following **macro** metrics:

Balanced Accuracy, F1, Sensitivity, Specificity, Positive Predictive Value, Negative Predictive Value, Kappa, Detection Rate, Detection Prevalence, and Prevalence.

In general, the metrics mentioned in binomial\_metrics() can be enabled as macro metrics (excluding MCC, AUC, Lower CI, Upper CI, and the AIC/AICc/BIC metrics). These metrics also has a weighted average version.

**N.B.** we also refer to the *one-vs-all evaluations* as the *class level results*.

#### **Multiclass metrics:**

In addition, the Overall Accuracy and *multiclass* MCC metrics are computed. *Multiclass* AUC can be enabled but is slow to calculate with many classes.

.....

#### The Summarized Results tibble contains:

Summary of the random evaluations.

**How**: The one-vs-all binomial evaluations are aggregated by repetition and summarized. Besides the metrics from the binomial evaluations, it also includes Overall Accuracy and *multiclass* MCC.

The Measure column indicates the statistical descriptor used on the evaluations. The Mean, Median, SD, IQR, Max, Min, NAs, and INFs measures describe the *Random Evaluations* tibble, while the CL\_Max, CL\_Min, CL\_NAs, and CL\_INFs describe the Class Level results.

The rows where Measure == All\_<<class name>> are the evaluations when all the observations are predicted to be in that class.

.....

#### The **Summarized Class Level Results** tibble contains:

The (nested) summarized results for each class, with the same metrics and descriptors as the *Summarized Results* tibble. Use tidyr::unnest on the tibble to inspect the results.

**How**: The one-vs-all evaluations are summarized by class.

The rows where Measure == All\_0 are the evaluations when none of the observations are predicted to be in that class, while the rows where Measure == All\_1 are the evaluations when all of the observations are predicted to be in that class.

.....

#### The **Random Evaluations** tibble contains:

The repetition results with the same metrics as the Summarized Results tibble.

**How**: The one-vs-all evaluations are aggregated by repetition. If a metric contains one or more NAs in the one-vs-all evaluations, it will lead to an NA result for that repetition.

Also includes:

A nested tibble with the one-vs-all binomial evaluations (**Class Level Results**), including nested **Confusion Matrices** and the **Support** column, which is a count of how many observations from the class is in the test set.

A nested tibble with the **predictions** and targets.

A list of **ROC** curve objects.

A nested tibble with the multiclass confusion matrix.

A nested **Process** information object with information about the evaluation.

Name of dependent variable.

#### Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

#### See Also

Other baseline functions: baseline(), baseline\_binomial(), baseline\_gaussian()

## **Examples**

```
# Attach packages
library(cvms)
library(groupdata2) # partition()
library(dplyr) # %>% arrange()
library(tibble)
# Data is part of cvms
data <- participant.scores</pre>
# Set seed for reproducibility
set.seed(1)
# Partition data
partitions <- partition(data, p = 0.7, list_out = TRUE)</pre>
train_set <- partitions[[1]]</pre>
test_set <- partitions[[2]]</pre>
# Create baseline evaluations
# Note: usually n=100 is a good setting
# Create some data with multiple classes
multiclass_data <- tibble(</pre>
  "target" = rep(paste0("class_", 1:5), each = 10)
  dplyr::sample_n(35)
baseline_multinomial(
  test_data = multiclass_data,
  dependent_col = "target",
  n = 4
# Parallelize evaluations
# Attach doParallel and register four cores
# Uncomment:
```

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```
# library(doParallel)
# registerDoParallel(4)
# Make sure to uncomment the parallel argument
(mb <- baseline_multinomial(</pre>
  test_data = multiclass_data,
  dependent_col = "target",
  n = 6
  #, parallel = TRUE # Uncomment
))
# Inspect the summarized class level results
# for class_2
mb$summarized_class_level_results %>%
  dplyr::filter(Class == "class_2") %>%
  tidyr::unnest(Results)
# Multinomial with custom random generator function
# that creates very "certain" predictions
# (once softmax is applied)
rcertain <- function(n) {</pre>
  (runif(n, min = 1, max = 100)^1.4) / 100
# Make sure to uncomment the parallel argument
baseline_multinomial(
  test_data = multiclass_data,
  dependent_col = "target",
  n = 6,
  random_generator_fn = rcertain
  #, parallel = TRUE # Uncomment
)
```

binomial\_metrics

Select metrics for binomial evaluation

## Description

#### [Experimental]

Enable/disable metrics for binomial evaluation. Can be supplied to the `metrics` argument in many of the cvms functions.

Note: Some functions may have slightly different defaults than the ones supplied here.

#### Usage

```
binomial_metrics(
   all = NULL,
```

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balanced\_accuracy = NULL,

accuracy = NULL,

```
f1 = NULL,
      sensitivity = NULL,
      specificity = NULL,
      pos_pred_value = NULL,
      neg_pred_value = NULL,
      auc = NULL,
      lower_ci = NULL,
      upper_ci = NULL,
      kappa = NULL,
     mcc = NULL,
      detection_rate = NULL,
      detection_prevalence = NULL,
      prevalence = NULL,
      false_neg_rate = NULL,
      false_pos_rate = NULL,
      false_discovery_rate = NULL,
      false_omission_rate = NULL,
      threat_score = NULL,
      aic = NULL,
      aicc = NULL,
     bic = NULL
   )
Arguments
   all
                    Enable/disable all arguments at once. (Logical)
                    Specifying other metrics will overwrite this, why you can use (all = FALSE,
                    accuracy = TRUE) to get only the Accuracy metric.
    balanced_accuracy
                    Balanced Accuracy (Default: TRUE)
                    Accuracy (Default: FALSE)
   accuracy
   f1
                    F1 (Default: TRUE)
                    Sensitivity (Default: TRUE)
   sensitivity
    specificity
                    Specificity (Default: TRUE)
   pos_pred_value Pos Pred Value (Default: TRUE)
   neg_pred_value Neg Pred Value (Default: TRUE)
                    AUC (Default: TRUE)
   auc
                    Lower CI (Default: TRUE)
    lower_ci
                    Upper CI (Default: TRUE)
   upper_ci
   kappa
                    Kappa (Default: TRUE)
                    MCC (Default: TRUE)
   mcc
```

detection\_rate Detection Rate (Default: TRUE)

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```
detection_prevalence
```

Detection Prevalence (Default: TRUE)

prevalence Prevalence (Default: TRUE)
false\_neg\_rate False Neg Rate (Default: FALSE)
false\_pos\_rate False Pos Rate (Default: FALSE)

false\_discovery\_rate

False Discovery Rate (Default: FALSE)

false\_omission\_rate

False Omission Rate (Default: FALSE)

aic AIC. (Default: FALSE)
aicc AICc. (Default: FALSE)
bic BIC. (Default: FALSE)

#### Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

#### See Also

Other evaluation functions: confusion\_matrix(), evaluate(), evaluate\_residuals(), gaussian\_metrics(), multinomial\_metrics()

#### **Examples**

```
# Attach packages
library(cvms)

# Enable only Balanced Accuracy
binomial_metrics(all = FALSE, balanced_accuracy = TRUE)

# Enable all but Balanced Accuracy
binomial_metrics(all = TRUE, balanced_accuracy = FALSE)

# Disable Balanced Accuracy
binomial_metrics(balanced_accuracy = FALSE)
```

combine\_predictors

Generate model formulas by combining predictors

#### **Description**

## [Maturing]

Create model formulas with every combination of your fixed effects, along with the dependent variable and random effects. 259,358 formulas have been precomputed with two- and three-way interactions for up to 8 fixed effects, with up to 5 included effects per formula. Uses the + and \* operators, so lower order interactions are automatically included.

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## Usage

```
combine_predictors(
  dependent,
  fixed_effects,
  random_effects = NULL,
  max_fixed_effects = 5,
  max_interaction_size = 3,
  max_effect_frequency = NULL
)
```

## **Arguments**

dependent Name of dependent variable. (Character)

fixed\_effects list of fixed effects. (Character)

Max. limit of 8 effects when interactions are included!

A fixed effect name cannot contain: white spaces, "\*" or "+".

Effects in sublists will be interchanged. This can be useful, when we have multiple versions of a predictor (e.g. x1 and log(x1)) that we do not wish to have

in the same formula.

Example of interchangeable effects:

list(list("x1", "log\_x1"), "x2", "x3")

random\_effects The random effects structure. (Character)

Is appended to the model formulas.

max\_fixed\_effects

Maximum number of fixed effects in a model formula. (Integer)

Max. limit of 5 when interactions are included!

max\_interaction\_size

Maximum number of effects in an interaction. (Integer)

Max. limit of 3.

Use this to limit the n-way interactions allowed. 0 or 1 excludes interactions all

A model formula can contain multiple interactions.

max\_effect\_frequency

Maximum number of times an effect is included in a formula string.

#### Value

list of model formulas.

```
E.g.:
```

```
c("y \sim x1 + (1|z)", "y \sim x2 + (1|z)", "y \sim x1 + x2 + (1|z)", "y \sim x1 * x2 + (1|z)").
```

## Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

#### **Examples**

```
# Attach packages
library(cvms)
# Create effect names
dependent <- "y"
fixed_effects <- c("a", "b", "c")
random_effects <- "(1|e)"</pre>
# Create model formulas
combine_predictors(
 dependent, fixed_effects,
 random_effects
)
# Create effect names with interchangeable effects in sublists
fixed_effects <- list("a", list("b", "log_b"), "c")</pre>
# Create model formulas
combine_predictors(
 dependent, fixed_effects,
 random_effects
)
```

compatible.formula.terms

Compatible formula terms

## **Description**

162,660 pairs of compatible terms for building model formulas with up to 15 fixed effects.

#### **Format**

```
A data. frame with 162,660 rows and 5 variables:
```

```
left term, fixed effect or interaction, with fixed effects separated by "*"
```

right term, fixed effect or interaction, with fixed effects separated by "\*"

max\_interaction\_size maximum interaction size in the two terms, up to 3

**num\_effects** number of unique fixed effects in the two terms, up to 5

min\_num\_fixed\_effects minimum number of fixed effects required to use a formula with the two terms, i.e. the index in the alphabet of the last of the alphabetically ordered effects (letters) in the two terms, so 4 if left == "A" and right == "D"

#### **Details**

A term is either a fixed effect or an interaction between fixed effects (up to three-way), where the effects are separated by the "\*" operator.

Two terms are compatible if they are not redundant, meaning that both add a fixed effect to the formula. E.g. as the interaction "x1 \* x2 \* x3" expands to "x1 + x2 + x3 + x1 \* x2 + x1 \* x3 + x2 \* x3 + x1 \* x2 \* x3", the higher order interaction makes these "sub terms" redundant. Note: All terms are compatible with NA.

Effects are represented by the first fifteen capital letters.

Used to generate the model formulas for combine\_predictors.

#### Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

confusion\_matrix

Create a confusion matrix

## **Description**

## [Experimental]

Creates a confusion matrix from targets and predictions. Calculates associated metrics.

Multiclass results are based on one-vs-all evaluations. Both regular averaging and weighted averaging are available. Also calculates the Overall Accuracy.

**Note**: In most cases you should use evaluate() instead. It has additional metrics and works in magrittr pipes (e.g. %>%) and with dplyr::group\_by(). confusion\_matrix() is more lightweight and may be preferred in programming when you don't need the extra stuff in evaluate().

#### Usage

```
confusion_matrix(
  targets,
  predictions,
  metrics = list(),
  positive = 2,
  c_levels = NULL,
  do_one_vs_all = TRUE,
  parallel = FALSE
)
```

## **Arguments**

```
targets vector with true classes. Either numeric or character.

predictions vector with predicted classes. Either numeric or character.
```

metrics

list for enabling/disabling metrics.

E.g. list("Accuracy" = TRUE) would add the regular accuracy metric, whie list("F1" = FALSE) would remove the F1 metric. Default values (TRUE/FALSE) will be used for the remaining available metrics.

You can enable/disable all metrics at once by including "all" = TRUE/FALSE in the list. This is done prior to enabling/disabling individual metrics, why for instance list("all" = FALSE, "Accuracy" = TRUE) would return only the Accuracy metric.

The list can be created with binomial\_metrics() or multinomial\_metrics(). Also accepts the string "all".

positive

Level from `targets` to predict. Either as character (*preferable*) or level index (1 or 2 - alphabetically). (**Two-class only**)

E.g. if we have the levels "cat" and "dog" and we want "dog" to be the positive class, we can either provide "dog" or 2, as alphabetically, "dog" comes after "cat".

**Note:** For *reproducibility*, it's preferable to **specify the name directly**, as different locales may sort the levels differently.

c\_levels

vector with categorical levels in the targets. Should have same type as `targets`.

If NULL, they are inferred from `targets`.

N.B. the levels are sorted alphabetically. When `positive` is numeric (i.e. an index), it therefore still refers to the index of the alphabetically sorted levels.

do\_one\_vs\_all

Whether to perform *one-vs-all* evaluations when working with more than 2

classes (multiclass).

If you are only interested in the confusion matrix, this allows you to skip most of the metric calculations.

parallel

Whether to perform the one-vs-all evaluations in parallel. (Logical)

N.B. This only makes sense when you have a lot of classes or a very large

dataset.

Remember to register a parallel backend first. E.g. with doParallel::registerDoParallel.

#### **Details**

The following formulas are used for calculating the metrics:

```
Sensitivity = TP / (TP + FN)
Specificity = TN / (TN + FP)
Pos Pred Value = TP / (TP + FP)
Neg Pred Value = TN / (TN + FN)
Balanced Accuracy = (Sensitivity + Specificity) / 2
Accuracy = (TP + TN) / (TP + TN + FP + FN)
Overall Accuracy = Correct / (Correct + Incorrect)
F1 = 2 * Pos Pred Value * Sensitivity / (Pos Pred Value + Sensitivity)
MCC = ((TP * TN) - (FP * FN)) / sqrt((TP + FP) * (TP + FN) * (TN + FP) * (TN + FN))
```

Note for MCC: Formula is for the *binary* case. When the denominator is 0, we set it to 1 to avoid NaN. See the metrics vignette for the multiclass version.

```
Detection Rate = TP / (TP + FN + TN + FP)
Detection Prevalence = (TP + FP) / (TP + FN + TN + FP)
Threat Score = TP / (TP + FN + FP)
False Neg Rate = 1 - Sensitivity
False Pos Rate = 1 - Specificity
False Discovery Rate = 1 - Pos Pred Value
False Omission Rate = 1 - Neg Pred Value
```

For **Kappa** the counts (TP, TN, FP, FN) are normalized to percentages (summing to 1). Then the following is calculated:

```
p_observed = TP + TN
p_expected = (TN + FP) * (TN + FN) + (FN + TP) * (FP + TP)
Kappa = (p_observed - p_expected) / (1 - p_expected)
```

#### Value

tibble with:

Nested **confusion matrix** (tidied version)

Nested confusion matrix (table)

The Positive Class.

Multiclass only: Nested **Class Level Results** with the two-class metrics, the nested confusion matrices, and the **Support** metric, which is a count of the class in the target column and is used for the weighted average metrics.

The following metrics are available (see `metrics`):

#### Two classes or more:

Metric	Name	Default
Balanced Accuracy	"Balanced Accuracy"	Enabled
Accuracy	"Accuracy"	Disabled
F1	"F1"	Enabled
Sensitivity	"Sensitivity"	Enabled
Specificity	"Specificity"	Enabled
Positive Predictive Value	"Pos Pred Value"	Enabled
Negative Predictive Value	"Neg Pred Value"	Enabled
Kappa	"Kappa"	Enabled
Matthews Correlation Coefficient	"MCC"	Enabled
Detection Rate	"Detection Rate"	Enabled
<b>Detection Prevalence</b>	"Detection Prevalence"	Enabled
Prevalence	"Prevalence"	Enabled
False Negative Rate	"False Neg Rate"	Disabled
False Positive Rate	"False Pos Rate"	Disabled
False Discovery Rate	"False Discovery Rate"	Disabled

False Omission Rate "False Omission Rate" Disabled
Threat Score "Threat Score" Disabled

The **Name** column refers to the name used in the package. This is the name in the output and when enabling/disabling in `metrics`.

#### Three classes or more:

The metrics mentioned above (excluding MCC) has a weighted average version (disabled by default; weighted by the **Support**).

In order to enable a weighted metric, prefix the metric name with "Weighted" when specifying `metrics`.

E.g. metrics = list("Weighted Accuracy" = TRUE).

Metric	Name	Default
Overall Accuracy	"Overall Accuracy"	Enabled
Weighted *	"Weighted *"	Disabled
Multiclass MCC	"MCC"	Enabled

## Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

#### See Also

Other evaluation functions: binomial\_metrics(), evaluate(), evaluate\_residuals(), gaussian\_metrics(), multinomial\_metrics()

## **Examples**

```
# Attach cvms
library(cvms)

# Two classes

# Create targets and predictions
targets <- c(0, 1, 0, 1, 0, 1, 0, 1, 0, 1, 0, 1)
predictions <- c(1, 1, 0, 0, 0, 1, 1, 1, 0, 1, 0, 0)

# Create confusion matrix with default metrics
cm <- confusion_matrix(targets, predictions)
cm
cm[["Confusion Matrix"]]
cm[["Table"]]

# Three classes

# Create targets and predictions
targets <- c(0, 1, 2, 1, 0, 1, 2, 1, 0, 1, 2, 1, 0)</pre>
```

```
predictions <- c(2, 1, 0, 2, 0, 1, 1, 2, 0, 1, 2, 0, 2)

# Create confusion matrix with default metrics
cm <- confusion_matrix(targets, predictions)
cm
cm[["Confusion Matrix"]]
cm[["Table"]]

# Enabling weighted accuracy

# Create confusion matrix with Weighted Accuracy enabled
cm <- confusion_matrix(targets, predictions,
    metrics = list("Weighted Accuracy" = TRUE)
)
cm</pre>
```

cross\_validate

Cross-validate regression models for model selection

## **Description**

#### [Stable]

Cross-validate one or multiple linear or logistic regression models at once. Perform repeated cross-validation. Returns results in a tibble for easy comparison, reporting and further analysis.

See cross\_validate\_fn() for use with custom model functions.

## Usage

```
cross_validate(
  data,
  formulas,
  family,
  fold_cols = ".folds",
  control = NULL,
  REML = FALSE,
  cutoff = 0.5,
  positive = 2,
  metrics = list(),
  preprocessing = NULL,
  rm_nc = FALSE,
  parallel = FALSE,
  verbose = FALSE,
  link = deprecated(),
 models = deprecated(),
  model_verbose = deprecated()
)
```

#### **Arguments**

data data.frame.

Must include one or more grouping factors for identifying folds - as made with

groupdata2::fold().

formulas Model formulas as strings. (Character)

> E.g. c("y~x", "y~z"). Can contain random effects.

E.g.  $c("y^x+(1|r)", "y^z+(1|r)")$ .

family Name of the family. (Character)

Currently supports "gaussian" for linear regression with lm() / lme4::lmer()

and "binomial" for binary classification with glm() / lme4::glmer().

See cross\_validate\_fn() for use with other model functions.

fold\_cols Name(s) of grouping factor(s) for identifying folds. (Character)

Include names of multiple grouping factors for repeated cross-validation.

Construct control structures for mixed model fitting (with lme4::lmer() or control

lme4::glmer()). See lme4::lmerControl and lme4::glmerControl.

N.B. Ignored if fitting lm() or glm() models.

REML Restricted Maximum Likelihood. (Logical)

cutoff Threshold for predicted classes. (Numeric)

N.B. Binomial models only

Level from dependent variable to predict. Either as character (preferable) or positive

level index (1 or 2 - alphabetically).

E.g. if we have the levels "cat" and "dog" and we want "dog" to be the positive class, we can either provide "dog" or 2, as alphabetically, "dog" comes after

Note: For reproducibility, it's preferable to specify the name directly, as different locales may sort the levels differently.

Used when calculating confusion matrix metrics and creating ROC curves.

The Process column in the output can be used to verify this setting.

N.B. Only affects evaluation metrics, not the model training or returned predic-

N.B. Binomial models only.

metrics list for enabling/disabling metrics.

E.g. list("RMSE" = FALSE) would remove RMSE from the results, and list("Accuracy"

= TRUE) would add the regular Accuracy metric to the classification results. Default values (TRUE/FALSE) will be used for the remaining available metrics.

You can enable/disable all metrics at once by including "all" = TRUE/FALSE in the list. This is done prior to enabling/disabling individual metrics, why list("all" = FALSE, "RMSE" = TRUE) would return only the RMSE metric.

The list can be created with gaussian\_metrics() or binomial\_metrics().

Also accepts the string "all".

preprocessing Name of preprocessing to apply.

Available preprocessings are:

Name "standardize' Centers and scales the numeric "range" Normalizes the numeric predictors to the 0-1 range. Values outside the min/max range in the test fold are trun "scale" Scales the numeric predictors to have a standard devia "center" Centers the numeric predictors to have a m The preprocessing parameters (mean, SD, etc.) are extracted from the training folds and applied to both the training folds and the test fold. They are returned in the **Preprocess** column for inspection. N.B. The preprocessings should not affect the results to a noticeable degree, although "range" might due to the truncation. rm\_nc Remove non-converged models from output. (Logical) parallel Whether to cross-validate the list of models in parallel. (Logical) Remember to register a parallel backend first. E.g. with doParallel::registerDoParallel. Whether to message process information like the number of model instances to verbose fit and which model function was applied. (Logical) link, models, model\_verbose

## Details

Packages used:

## **Models:**

Gaussian: stats::lm, lme4::lmer
Binomial: stats::glm, lme4::glmer

Deprecated.

## **Results:**

Shared:

AIC: stats::AIC
AICc: MuMIn::AICc
BIC: stats::BIC

Gaussian:

r2m: MuMIn::r.squaredGLMM
r2c: MuMIn::r.squaredGLMM

Binomial:

ROC and AUC: pROC::roc

#### Value

tibble with results for each model.

**Shared across families:** A nested tibble with **coefficients** of the models from all iterations.

Number of *total* **folds**.

Number of **fold columns**.

Count of **convergence warnings**. Consider discarding models that did not converge on all iterations. Note: you might still see results, but these should be taken with a grain of salt!

Count of other warnings. These are warnings without keywords such as "convergence".

Count of Singular Fit messages. See lme4::isSingular for more information.

Nested tibble with the warnings and messages caught for each model.

A nested **Process** information object with information about the evaluation.

Name of **dependent** variable.

Names of fixed effects.

Names of random effects, if any.

Nested tibble with **preprocess**ing parameters, if any.

## **Gaussian Results:**

\_\_\_\_\_

Average RMSE, MAE, NRMSE(IQR), RRSE, RAE, RMSLE, AIC, AICc, and BIC of all the iterations\*, **omitting potential NAs** *from non-converged iterations*. Note that the Information Criterion metrics (AIC, AICc, and BIC) are also averages.

See the additional metrics (disabled by default) at ?gaussian\_metrics.

A nested tibble with the **predictions** and targets.

A nested tibble with the non-averaged results from all iterations.

\* In *repeated cross-validation*, the metrics are first averaged for each fold column (repetition) and then averaged again.

#### **Binomial Results:**

Based on the **collected** predictions from the test folds\*, a confusion matrix and a ROC curve are created to get the following:

ROC:

AUC, Lower CI, and Upper CI

 ${\tt Confusion\,Matrix:}$ 

Balanced Accuracy, F1, Sensitivity, Specificity, Positive Predictive Value, Negative Predictive Value, Kappa, Detection Rate, Detection Prevalence, Prevalence, and MCC (Matthews correlation coefficient).

See the additional metrics (disabled by default) at ?binomial\_metrics.

Also includes:

A nested tibble with **predictions**, predicted classes (depends on cutoff), and the targets. Note, that the predictions are *not necessarily* of the *specified* positive class, but of the *model's* positive class (second level of dependent variable, alphabetically).

```
The pROC::roc ROC curve object(s).
```

A nested tibble with the **confusion matrix**/matrices. The Pos\_ columns tells you whether a row is a True Positive (TP), True Negative (TN), False Positive (FP), or False Negative (FN), depending on which level is the "positive" class. I.e. the level you wish to predict.

A nested tibble with the **results** from all fold columns.

The name of the **Positive Class**.

\* In repeated cross-validation, an evaluation is made per fold column (repetition) and averaged.

#### Author(s)

```
Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>
Benjamin Hugh Zachariae
```

#### See Also

Other validation functions: cross\_validate\_fn(), validate(), validate\_fn()

## **Examples**

```
# Attach packages
library(cvms)
library(groupdata2) # fold()
library(dplyr) # %>% arrange()
# Data is part of cvms
data <- participant.scores</pre>
# Set seed for reproducibility
set.seed(7)
# Fold data
data <- fold(</pre>
  data,
  k = 4
  cat_col = "diagnosis",
  id_col = "participant"
  arrange(.folds)
# Cross-validate a single model
# Gaussian
cross_validate(
  data,
  formulas = "score~diagnosis",
```

```
family = "gaussian",
  REML = FALSE
)
# Binomial
cross_validate(
  data,
  formulas = "diagnosis~score",
  family = "binomial"
)
# Cross-validate multiple models
formulas <- c(</pre>
  "score~diagnosis+(1|session)",
  "score~age+(1|session)"
)
cross_validate(
  data,
  formulas = formulas,
  family = "gaussian",
  REML = FALSE
)
# Use parallelization
# Attach doParallel and register four cores
# Uncomment:
# library(doParallel)
# registerDoParallel(4)
# Cross-validate a list of model formulas in parallel
# Make sure to uncomment the parallel argument
cross_validate(
  data,
  formulas = formulas,
  family = "gaussian"
  #, parallel = TRUE # Uncomment
)
```

## **Description**

#### [Experimental]

Cross-validate your model function with one or multiple model formulas at once. Perform repeated cross-validation. Preprocess the train/test split within the cross-validation. Perform hyperparameter tuning with grid search. Returns results in a tibble for easy comparison, reporting and further analysis.

Compared to cross\_validate(), this function allows you supply a custom model function, a predict function, a preprocess function and the hyperparameter values to cross-validate.

Supports regression and classification (binary and multiclass). See `type`.

Note that some metrics may not be computable for some types of model objects.

## Usage

```
cross_validate_fn(
  data,
  formulas,
  type,
  model_fn,
  predict_fn,
  preprocess_fn = NULL,
  preprocess_once = FALSE,
  hyperparameters = NULL,
  fold_cols = ".folds",
  cutoff = 0.5,
  positive = 2,
  metrics = list(),
  rm_nc = FALSE,
  parallel = FALSE,
  verbose = TRUE
)
```

#### **Arguments**

```
data data.frame.

Must include one or more grouping factors for identifying folds - as made with groupdata2::fold().

formulas

Model formulas as strings. (Character)

Will be converted to formula objects before being passed to `model_fn`.

E.g. c("y~x", "y~z").

Can contain random effects.

E.g. c("y~x+(1|r)", "y~z+(1|r)").

type

Type of evaluation to perform:

"gaussian" for regression (like linear regression).

"binomial" for binary classification.

"multinomial" for multiclass classification.
```

model\_fn

Model function that returns a fitted model object. Will usually wrap an existing model function like e1071::svm or nnet::multinom.

Must have the following function arguments:

predict\_fn

Function for predicting the targets in the test folds/sets using the fitted model object. Will usually wrap stats::predict(), but doesn't have to.

Must have the following function arguments:

Must return predictions in the following formats, depending on `type`:

**Binomial:** vector or one-column matrix / data.frame with probabilities (0-1) of the second class, alphabetically. E.g.:

```
c(0.3, 0.5, 0.1, 0.5)
```

N.B. When unsure whether a model type produces probabilities based off the alphabetic order of your classes, using 0 and 1 as classes in the dependent variable instead of the class names should increase the chance of getting probabilities of the right class.

**Gaussian:** vector or one-column matrix / data.frame with the predicted value. E.g.:

```
c(3.7, 0.9, 1.2, 7.3)
```

**Multinomial:** data. frame with one column per class containing probabilities of the class. Column names should be identical to how the class names are written in the target column. E.g.:

class_1	class_2	class_3
0.269	0.528	0.203
0.368	0.322	0.310
0.375	0.371	0.254

preprocess\_fn Function for preprocessing the training and test sets.

Can, for instance, be used to standardize both the training and test sets with the scaling and centering parameters from the training set.

Must have the following function arguments:

Must return a list with the preprocessed `train\_data` and `test\_data`. It may also contain a tibble with the parameters used in preprocessing:

```
list("train" = train_data,
     "test" = test_data,
     "parameters" = preprocess_parameters)
```

Additional elements in the returned list will be ignored.

The optional parameters tibble will be included in the output. It could have the following format:

Measure	var_1	var_2
Mean	37.921	88.231
SD	12.4	5.986

N.B. When `preprocess\_once` is FALSE, the current formula and hyperparameters will be provided. Otherwise, these arguments will be NULL.

#### preprocess\_once

Whether to apply the preprocessing once (**ignoring** the formula and hyperparameters arguments in `preprocess\_fn`) or for every model separately. (Logical)

When preprocessing does not depend on the current formula or hyperparameters, we can do the preprocessing of each train/test split once, to save time. This may require holding a lot more data in memory though, why it is not the default setting.

#### hyperparameters

Either a named list with hyperparameter values to combine in a grid or a data.frame with one row per hyperparameter combination.

Named list for grid search: Add ".n" to sample the combinations. Can be the number of combinations to use, or a percentage between  $\emptyset$  and 1.

```
list(".n" = 10, # sample 10 combinations
    "lrn_rate" = c(0.1, 0.01, 0.001),
    "h_layers" = c(10, 100, 1000),
    "drop_out" = runif(5, 0.3, 0.7))
```

data.frame with specific hyperparameter combinations: One row per combination to test.

E.g.

lrn_rate	h_layers	drop_out
0.1	10	0.65
0.1	1000	0.65
0.01	1000	0.63

fold\_cols

Name(s) of grouping factor(s) for identifying folds. (Character)

Include names of multiple grouping factors for repeated cross-validation.

cutoff

Threshold for predicted classes. (Numeric)

#### N.B. Binomial models only

positive

Level from dependent variable to predict. Either as character (*preferable*) or level index (1 or 2 - alphabetically).

E.g. if we have the levels "cat" and "dog" and we want "dog" to be the positive class, we can either provide "dog" or 2, as alphabetically, "dog" comes after "cat".

**Note:** For *reproducibility*, it's preferable to **specify the name directly**, as different locales may sort the levels differently.

Used when calculating confusion matrix metrics and creating ROC curves.

The Process column in the output can be used to verify this setting.

N.B. Only affects evaluation metrics, not the model training or returned predictions.

### N.B. Binomial models only.

metrics

list for enabling/disabling metrics.

E.g. list("RMSE" = FALSE) would remove RMSE from the regression results, and list("Accuracy" = TRUE) would add the regular Accuracy metric to the classification results. Default values (TRUE/FALSE) will be used for the remaining available metrics.

You can enable/disable all metrics at once by including "all" = TRUE/FALSE in the list. This is done prior to enabling/disabling individual metrics, why f.i. list("all" = FALSE, "RMSE" = TRUE) would return only the RMSE metric.

The list can be created with gaussian\_metrics(), binomial\_metrics(), or multinomial\_metrics().

Also accepts the string "all".

rm\_nc Remove non-converged models from output. (Logical)

parallel Whether to cross-validate the list of models in parallel. (Logical)

Remember to register a parallel backend first. E.g. with doParallel::registerDoParallel.

verbose Whether to message process information like the number of model instances to

fit. (Logical)

#### **Details**

## Packages used:

## **Results:**

Shared:

AIC: stats::AIC AICc: MuMIn::AICc BIC: stats::BIC

Gaussian:

r2m: MuMIn::r.squaredGLMM r2c: MuMIn::r.squaredGLMM Binomial and Multinomial:

ROC and related metrics: Binomial: pROC::roc

 $Multinomial: \verb"pROC::multiclass.roc"$ 

# Value

tibble with results for each model.

N.B. The **Fold** column in the nested tibbles contains the test fold in that train/test split.

#### **Shared across families:**

A nested tibble with **coefficients** of the models from all iterations. The coefficients are extracted from the model object with parameters::model\_parameters() or coef() (with some restrictions on the output). If these attempts fail, a default coefficients tibble filled with NAs is returned.

Nested tibble with the used **preprocessing parameters**, if a passed preprocess\_fn returns the parameters in a tibble.

Number of total folds.

Number of **fold columns**.

Count of **convergence warnings**, using a limited set of keywords (e.g. "convergence"). If a convergence warning does not contain one of these keywords, it will be counted with **other warnings**. Consider discarding models that did not converge on all iterations. Note: you might still see results, but these should be taken with a grain of salt!

Nested tibble with the warnings and messages caught for each model.

A nested **Process** information object with information about the evaluation.

Name of **dependent** variable.

Names of **fixed** effects.

Names of random effects, if any.

\_\_\_\_\_

# **Gaussian Results:**

Average RMSE, MAE, NRMSE (IQR), RRSE, RAE, RMSLE of all the iterations\*, **omitting potential NAs** *from non-converged iterations*.

See the additional metrics (disabled by default) at ?gaussian\_metrics.

A nested tibble with the **predictions** and targets.

A nested tibble with the non-averaged **results** from all iterations.

\* In *repeated cross-validation*, the metrics are first averaged for each fold column (repetition) and then averaged again.

# **Binomial Results:**

Based on the **collected** predictions from the test folds\*, a confusion matrix and a ROC curve are created to get the following:

ROC:

AUC, Lower CI, and Upper CI

Confusion Matrix:

Balanced Accuracy, F1, Sensitivity, Specificity, Positive Predictive Value, Negative Predictive Value, Kappa, Detection Rate, Detection Prevalence, Prevalence, and MCC (Matthews correlation coefficient).

See the additional metrics (disabled by default) at ?binomial\_metrics.

Also includes:

A nested tibble with **predictions**, predicted classes (depends on cutoff), and the targets. Note, that the predictions are *not necessarily* of the *specified* positive class, but of the *model's* positive class (second level of dependent variable, alphabetically).

The pROC::roc ROC curve object(s).

A nested tibble with the **confusion matrix**/matrices. The Pos\_ columns tells you whether a row is a True Positive (TP), True Negative (TN), False Positive (FP), or False Negative (FN), depending on which level is the "positive" class. I.e. the level you wish to predict.

A nested tibble with the **results** from all fold columns.

The name of the **Positive Class**.

\* In repeated cross-validation, an evaluation is made per fold column (repetition) and averaged.

# **Multinomial Results:**

For each class, a *one-vs-all* binomial evaluation is performed. This creates a **Class Level Results** tibble containing the same metrics as the binomial results described above (excluding MCC, AUC, Lower CI and Upper CI), along with a count of the class in the target column (Support). These metrics are used to calculate the **macro-averaged** metrics. The nested class level results tibble is also included in the output tibble, and could be reported along with the macro and overall metrics.

The output tibble contains the macro and overall metrics. The metrics that share their name with the metrics in the nested class level results tibble are averages of those metrics (note: does not remove NAs before averaging). In addition to these, it also includes the Overall Accuracy and the multiclass MCC.

**Note:** Balanced Accuracy is the macro-averaged metric, *not* the macro sensitivity as sometimes used!

Other available metrics (disabled by default, see metrics): Accuracy, *multiclass* AUC, Weighted Balanced Accuracy, Weighted Accuracy, Weighted F1, Weighted Sensitivity, Weighted Sensitivity, Weighted Sensitivity, Weighted Pos Pred Value, Weighted Neg Pred Value, Weighted Kappa, Weighted Detection Rate, Weighted Detection Prevalence, and Weighted Prevalence.

Note that the "Weighted" average metrics are weighted by the Support.

Also includes:

A nested tibble with the **predictions**, predicted classes, and targets.

A list of **ROC** curve objects when AUC is enabled.

A nested tibble with the multiclass **Confusion Matrix**.

# **Class Level Results**

Besides the binomial evaluation metrics and the Support, the nested class level results tibble also contains a nested tibble with the **Confusion Matrix** from the one-vs-all evaluation. The Pos\_ columns tells you whether a row is a True Positive (TP), True Negative (TN), False Positive (FP), or False Negative (FN), depending on which level is the "positive" class. In our case, 1 is the current class and 0 represents all the other classes together.

## Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

### See Also

Other validation functions: cross\_validate(), validate(), validate\_fn()

# **Examples**

```
# Attach packages
library(cvms)
library(groupdata2) # fold()
library(dplyr) # %>% arrange() mutate()
# Note: More examples of custom functions can be found at:
# model_fn: model_functions()
# predict_fn: predict_functions()
# preprocess_fn: preprocess_functions()
# Data is part of cvms
data <- participant.scores</pre>
# Set seed for reproducibility
set.seed(7)
# Fold data
data <- fold(</pre>
  data,
  k = 4,
 cat_col = "diagnosis",
  id_col = "participant"
  mutate(diagnosis = as.factor(diagnosis)) %>%
  arrange(.folds)
# Cross-validate multiple formulas
formulas_gaussian <- c(</pre>
  "score ~ diagnosis",
  "score ~ age"
)
formulas_binomial <- c(</pre>
  "diagnosis ~ score",
  "diagnosis ~ age"
)
# Gaussian
# Create model function that returns a fitted model object
lm_model_fn <- function(train_data, formula, hyperparameters) {</pre>
  lm(formula = formula, data = train_data)
}
# Create predict function that returns the predictions
```

```
lm_predict_fn <- function(test_data, model, formula,</pre>
                          hyperparameters, train_data) {
 stats::predict(
   object = model,
   newdata = test_data,
   type = "response",
   allow.new.levels = TRUE
 )
}
# Cross-validate the model function
cross_validate_fn(
 data,
 formulas = formulas_gaussian,
 type = "gaussian",
 model_fn = lm_model_fn,
 predict_fn = lm_predict_fn,
 fold_cols = ".folds"
)
# Binomial
# Create model function that returns a fitted model object
glm_model_fn <- function(train_data, formula, hyperparameters) {</pre>
 glm(formula = formula, data = train_data, family = "binomial")
# Create predict function that returns the predictions
glm_predict_fn <- function(test_data, model, formula,</pre>
                           hyperparameters, train_data) {
 stats::predict(
   object = model,
   newdata = test_data,
   type = "response",
   allow.new.levels = TRUE
 )
}
# Cross-validate the model function
cross_validate_fn(
 data,
 formulas = formulas_binomial,
 type = "binomial",
 model_fn = glm_model_fn,
 predict_fn = glm_predict_fn,
 fold_cols = ".folds"
)
# Support Vector Machine (svm)
# with hyperparameter tuning
```

```
# Only run if the `e1071` package is installed
if (requireNamespace("e1071", quietly = TRUE)){
# Create model function that returns a fitted model object
# We use the hyperparameters arg to pass in the kernel and cost values
svm_model_fn <- function(train_data, formula, hyperparameters) {</pre>
 # Expected hyperparameters:
 # - kernel
 # - cost
 if (!"kernel" %in% names(hyperparameters))
    stop("'hyperparameters' must include 'kernel'")
 if (!"cost" %in% names(hyperparameters))
    stop("'hyperparameters' must include 'cost'")
 e1071::svm(
    formula = formula,
   data = train_data,
   kernel = hyperparameters[["kernel"]],
   cost = hyperparameters[["cost"]],
   scale = FALSE,
   type = "C-classification",
   probability = TRUE
 )
}
# Create predict function that returns the predictions
svm_predict_fn <- function(test_data, model, formula,</pre>
                           hyperparameters, train_data) {
 predictions <- stats::predict(</pre>
   object = model,
   newdata = test_data,
   allow.new.levels = TRUE,
   probability = TRUE
 )
 # Extract probabilities
 probabilities <- dplyr::as_tibble(</pre>
   attr(predictions, "probabilities")
 # Return second column
 probabilities[[2]]
}
# Specify hyperparameters to try
# The optional ".n" samples 4 combinations
svm_hparams <- list(</pre>
 ".n" = 4,
  "kernel" = c("linear", "radial"),
  "cost" = c(1, 5, 10)
```

```
# Cross-validate the model function
cv <- cross_validate_fn(</pre>
  data,
  formulas = formulas_binomial,
  type = "binomial",
  model_fn = svm_model_fn,
  predict_fn = svm_predict_fn,
  hyperparameters = svm_hparams,
  fold_cols = ".folds"
)
C۷
# The `HParams` column has the nested hyperparameter values
cv %>%
  select(Dependent, Fixed, HParams, `Balanced Accuracy`, F1, AUC, MCC) %>%
  tidyr::unnest(cols = "HParams") %>%
  arrange(desc(`Balanced Accuracy`), desc(F1))
# Use parallelization
# The below examples show the speed gains when running in parallel
# Attach doParallel and register four cores
# Uncomment:
# library(doParallel)
# registerDoParallel(4)
# Specify hyperparameters such that we will
# cross-validate 20 models
hparams <- list(</pre>
  "kernel" = c("linear", "radial"),
  "cost" = 1:5
# Cross-validate a list of 20 models in parallel
# Make sure to uncomment the parallel argument
system.time({
  cross_validate_fn(
    data,
    formulas = formulas_gaussian,
    type = "gaussian",
    model_fn = svm_model_fn,
   predict_fn = svm_predict_fn,
   hyperparameters = hparams,
    fold_cols = ".folds"
    #, parallel = TRUE # Uncomment
  )
})
```

```
# Cross-validate a list of 20 models sequentially
system.time({
   cross_validate_fn(
        data,
        formulas = formulas_gaussian,
        type = "gaussian",
        model_fn = svm_model_fn,
        predict_fn = svm_predict_fn,
        hyperparameters = hparams,
        fold_cols = ".folds"
        #, parallel = TRUE # Uncomment
)
} # closes `e1071` package check
```

evaluate

Evaluate your model's performance

# Description

# [Maturing]

Evaluate your model's predictions on a set of evaluation metrics.

Create ID-aggregated evaluations by multiple methods.

Currently supports regression and classification (binary and multiclass). See `type`.

# Usage

```
evaluate(
   data,
   target_col,
   prediction_cols,
   type,
   id_col = NULL,
   id_method = "mean",
   apply_softmax = FALSE,
   cutoff = 0.5,
   positive = 2,
   metrics = list(),
   include_predictions = TRUE,
   parallel = FALSE,
   models = deprecated()
)
```

# **Arguments**

data

data.frame with predictions, targets and (optionally) an ID column. Can be grouped with group\_by.

**Multinomial:** When `type` is "multinomial", the predictions can be passed in one of two formats.

*Probabilities (Preferable):* 

One column per class with the probability of that class. The columns should have the name of their class, as they are named in the target column. E.g.:

target	class_3	class_2	class_1
class_2	0.203	0.528	0.269
class_3	0.310	0.322	0.368
class_2	0.254	0.371	0.375

### Classes:

A single column of type character with the predicted classes. E.g.:

prediction	target
class_2	class_2
class_1	class_3
class_1	class_2

**Binomial:** When `type` is "binomial", the predictions can be passed in one of two formats.

*Probabilities (Preferable):* One column with the **probability of class being the second class alphabetically** (1 if classes are 0 and 1). E.g.:

target	prediction
1	0.769
1	0.368
0	0.375

Note: At the alphabetical ordering of the class labels, they are of type character, why e.g. 100 would come before 7.

# Classes:

A single column of type character with the predicted classes. E.g.:

prediction	target
class_0	class_1
class_1	class_1
class_1	class_0

Note: The prediction column will be converted to the probability 0.0 for the first class alphabetically and 1.0 for the second class alphabetically.

> Gaussian: When `type` is "gaussian", the predictions should be passed as one column with the predicted values. E.g.:

prediction	target
28.9	30.2
33.2	27.1
23.4	21.3
•••	

Name of the column with the true classes/values in `data`. target\_col

> When `type` is "multinomial", this column should contain the class names, not their indices.

prediction\_cols

Name(s) of column(s) with the predictions.

Columns can be either numeric or character depending on which format is chosen. See 'data' for the possible formats.

Type of evaluation to perform:

"gaussian" for regression (like linear regression).

"binomial" for binary classification.

"multinomial" for multiclass classification.

id\_col Name of ID column to aggregate predictions by.

> N.B. Current methods assume that the target class/value is constant within the IDs.

N.B. When aggregating by ID, some metrics may be disabled.

id\_method Method to use when aggregating predictions by ID. Either "mean" or "majority".

When 'type' is gaussian, only the "mean" method is available.

mean: The average prediction (value or probability) is calculated per ID and evaluated. This method assumes that the target class/value is constant within the IDs.

majority: The most predicted class per ID is found and evaluated. In case of a tie, the winning classes share the probability (e.g. P = 0.5 each when two majority classes). This method assumes that the target class/value is constant within the IDs.

apply\_softmax Whether to apply the softmax function to the prediction columns when `type`

is "multinomial".

N.B. Multinomial models only.

cutoff Threshold for predicted classes. (Numeric)

N.B. Binomial models only.

Level from dependent variable to predict. Either as character (preferable) or

level index (1 or 2 - alphabetically).

E.g. if we have the levels "cat" and "dog" and we want "dog" to be the positive class, we can either provide "dog" or 2, as alphabetically, "dog" comes after "cat".

Note: For reproducibility, it's preferable to specify the name directly, as different locales may sort the levels differently.

type

positive

Used when calculating confusion matrix metrics and creating ROC curves.

The Process column in the output can be used to verify this setting.

N.B. Only affects the evaluation metrics. **Does NOT affect what the probabilities are of (always the second class alphabetically).** 

N.B. Binomial models only.

metrics

list for enabling/disabling metrics.

E.g. list("RMSE" = FALSE) would remove RMSE from the regression results, and list("Accuracy" = TRUE) would add the regular Accuracy metric to the classification results. Default values (TRUE/FALSE) will be used for the remaining available metrics.

You can enable/disable all metrics at once by including "all" = TRUE/FALSE in the list. This is done prior to enabling/disabling individual metrics, why f.i. list("all" = FALSE, "RMSE" = TRUE) would return only the RMSE metric.

The list can be created with gaussian\_metrics(), binomial\_metrics(), or multinomial\_metrics().

Also accepts the string "all".

include\_predictions

Whether to include the predictions in the output as a nested tibble. (Logical)

parallel Whether to run evaluations in parallel, when `data` is grouped with group\_by.

models Deprecated.

#### **Details**

Packages used:

Binomial and Multinomial:

ROC and AUC:

Binomial: pROC::roc

Multinomial: pROC::multiclass.roc

## Value

#### Gaussian Results:

tibble containing the following metrics by default:

Average RMSE, MAE, NRMSE(IQR), RRSE, RAE, RMSLE.

See the additional metrics (disabled by default) at ?gaussian\_metrics.

Also includes:

A nested tibble with the **Predictions** and targets.

A nested **Process** information object with information about the evaluation.

### **Binomial Results:**

\_\_\_\_\_\_

tibble with the following evaluation metrics, based on a confusion matrix and a ROC curve fitted to the predictions:

Confusion Matrix:

Balanced Accuracy, Accuracy, F1, Sensitivity, Specificity, Positive Predictive Value, Negative Predictive Value, Kappa, Detection Rate, Detection Prevalence, Prevalence, and MCC (Matthews correlation coefficient).

ROC:

AUC, Lower CI, and Upper CI

Note, that the ROC curve is only computed if AUC is enabled. See metrics.

Also includes:

A nested tibble with the **predictions** and targets.

A list of **ROC** curve objects (if computed).

A nested tibble with the **confusion matrix**. The Pos\_ columns tells you whether a row is a True Positive (TP), True Negative (TN), False Positive (FP), or False Negative (FN), depending on which level is the "positive" class. I.e. the level you wish to predict.

A nested **Process** information object with information about the evaluation.

### **Multinomial Results:**

For each class, a *one-vs-all* binomial evaluation is performed. This creates a **Class Level Results** tibble containing the same metrics as the binomial results described above (excluding Accuracy, MCC, AUC, Lower CI and Upper CI), along with a count of the class in the target column (Support). These metrics are used to calculate the **macro-averaged** metrics. The nested class level results tibble is also included in the output tibble, and could be reported along with the macro and overall metrics.

The output tibble contains the macro and overall metrics. The metrics that share their name with the metrics in the nested class level results tibble are averages of those metrics (note: does not remove NAs before averaging). In addition to these, it also includes the Overall Accuracy and the multiclass MCC.

**Note:** Balanced Accuracy is the macro-averaged metric, *not* the macro sensitivity as sometimes used!

Other available metrics (disabled by default, see metrics): Accuracy, *multiclass* AUC, Weighted Balanced Accuracy, Weighted Accuracy, Weighted F1, Weighted Sensitivity, Weighted Sensitivity, Weighted Specificity, Weighted Pos Pred Value, Weighted Neg Pred Value, Weighted Kappa, Weighted Detection Rate, Weighted Detection Prevalence, and Weighted Prevalence.

Note that the "Weighted" average metrics are weighted by the Support.

When having a large set of classes, consider keeping AUC disabled.

Also includes:

A nested tibble with the **Predictions** and targets.

A list of **ROC** curve objects when AUC is enabled.

A nested tibble with the multiclass Confusion Matrix.

A nested **Process** information object with information about the evaluation.

#### Class Level Results:

Besides the binomial evaluation metrics and the Support, the nested class level results tibble also contains a nested tibble with the **Confusion Matrix** from the one-vs-all evaluation. The Pos\_ columns tells you whether a row is a True Positive (TP), True Negative (TN), False Positive (FP), or False Negative (FN), depending on which level is the "positive" class. In our case, 1 is the current class and 0 represents all the other classes together.

#### Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

#### See Also

```
Other evaluation functions: binomial_metrics(), confusion_matrix(), evaluate_residuals(), gaussian_metrics(), multinomial_metrics()
```

## **Examples**

```
# Attach packages
library(cvms)
library(dplyr)
# Load data
data <- participant.scores
# Fit models
gaussian_model <- lm(age ~ diagnosis, data = data)</pre>
binomial_model <- glm(diagnosis ~ score, data = data)</pre>
# Add predictions
data[["gaussian_predictions"]] <- predict(gaussian_model, data,</pre>
  type = "response",
  allow.new.levels = TRUE
data[["binomial_predictions"]] <- predict(binomial_model, data,</pre>
  allow.new.levels = TRUE
# Gaussian evaluation
evaluate(
  data = data, target_col = "age",
  prediction_cols = "gaussian_predictions",
  type = "gaussian"
)
# Binomial evaluation
evaluate(
  data = data, target_col = "diagnosis",
  prediction_cols = "binomial_predictions",
  type = "binomial"
)
```

```
# Multinomial
# Create a tibble with predicted probabilities and targets
data_mc <- multiclass_probability_tibble(</pre>
  num_classes = 3, num_observations = 45,
  apply_softmax = TRUE, FUN = runif,
  class_name = "class_",
  add_targets = TRUE
)
class_names <- paste0("class_", 1:3)</pre>
# Multinomial evaluation
evaluate(
  data = data_mc, target_col = "Target",
  prediction_cols = class_names,
  type = "multinomial"
)
# ID evaluation
# Gaussian ID evaluation
# Note that 'age' is the same for all observations
# of a participant
evaluate(
  data = data, target_col = "age",
  prediction_cols = "gaussian_predictions",
  id_col = "participant",
  type = "gaussian"
)
# Binomial ID evaluation
evaluate(
  data = data, target_col = "diagnosis",
  prediction_cols = "binomial_predictions",
  id_col = "participant",
  id_method = "mean", # alternatively: "majority"
  type = "binomial"
)
# Multinomial ID evaluation
# Add IDs and new targets (must be constant within IDs)
data_mc[["Target"]] <- NULL</pre>
data_mc[["ID"]] <- rep(1:9, each = 5)</pre>
id_classes <- tibble::tibble(</pre>
  "ID" = 1:9,
  "Target" = sample(x = class_names, size = 9, replace = TRUE)
)
```

```
data_mc <- data_mc %>%
 dplyr::left_join(id_classes, by = "ID")
# Perform ID evaluation
evaluate(
 data = data_mc, target_col = "Target",
 prediction_cols = class_names,
 id_col = "ID",
 id_method = "mean", # alternatively: "majority"
 type = "multinomial"
)
# Training and evaluating a multinomial model with nnet
# Only run if `nnet` is installed
if (requireNamespace("nnet", quietly = TRUE)){
# Create a data frame with some predictors and a target column
class_names <- paste0("class_", 1:4)</pre>
data_for_nnet <- multiclass_probability_tibble(</pre>
 num_classes = 3, # Here, number of predictors
 num_observations = 30,
 apply_softmax = FALSE,
 FUN = rnorm,
 class_name = "predictor_"
 dplyr::mutate(Target = sample(
   class_names,
   size = 30,
    replace = TRUE
 ))
# Train multinomial model using the nnet package
mn_model <- nnet::multinom(</pre>
  "Target ~ predictor_1 + predictor_2 + predictor_3",
 data = data_for_nnet
# Predict the targets in the dataset
# (we would usually use a test set instead)
predictions <- predict(</pre>
 mn_model,
 data_for_nnet,
 type = "probs"
) %>%
 dplyr::as_tibble()
# Add the targets
predictions[["Target"]] <- data_for_nnet[["Target"]]</pre>
# Evaluate predictions
```

54 evaluate\_residuals

```
evaluate(
  data = predictions,
  target_col = "Target",
  prediction_cols = class_names,
  type = "multinomial"
)
}
```

evaluate\_residuals

Evaluate residuals from a regression task

# **Description**

# [Experimental]

Calculates a large set of error metrics from regression residuals.

**Note**: In most cases you should use evaluate() instead. It works in magrittr pipes (e.g. %>%) and with dplyr::group\_by(). evaluate\_residuals() is more lightweight and may be preferred in programming when you don't need the extra stuff in evaluate().

# Usage

```
evaluate_residuals(data, target_col, prediction_col, metrics = list())
```

### **Arguments**

data data. frame with predictions and targets.

target\_col Name of the column with the true values in `data`.

prediction\_col Name of column with the predicted values in `data`.

metrics list for enabling/disabling metrics.

 $E.g.\ \texttt{list("RMSE"} = \texttt{FALSE})\ would\ disable\ \texttt{RMSE}.\ Default\ values\ (\texttt{TRUE/FALSE})$ 

will be used for the remaining available metrics.

You can enable/disable all metrics at once by including "all" = TRUE/FALSE in the list. This is done prior to enabling/disabling individual metrics, why for instance list("all" = FALSE, "RMSE" = TRUE) would return only the RMSE

metric.

The list can be created with gaussian\_metrics().

Also accepts the string "all".

#### **Details**

The metric formulas are listed in 'The Available Metrics' vignette.

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# Value

tibble data. frame with the calculated metrics.

The following metrics are available (see `metrics`):

Metric	Name	Default
Mean Absolute Error	"MAE"	Enabled
Root Mean Square Error	"RMSE"	Enabled
Normalized RMSE (by target range)	"NRMSE(RNG)"	Disabled
Normalized RMSE (by target IQR)	"NRMSE(IQR)"	Enabled
Normalized RMSE (by target STD)	"NRMSE(STD)"	Disabled
Normalized RMSE (by target mean)	"NRMSE(AVG)"	Disabled
Relative Squared Error	"RSE"	Disabled
Root Relative Squared Error	"RRSE"	Enabled
Relative Absolute Error	"RAE"	Enabled
Root Mean Squared Log Error	"RMSLE"	Enabled
Mean Absolute Log Error	"MALE"	Disabled
Mean Absolute Percentage Error	"MAPE"	Disabled
Mean Squared Error	"MSE"	Disabled
Total Absolute Error	"TAE"	Disabled
Total Squared Error	"TSE"	Disabled

The **Name** column refers to the name used in the package. This is the name in the output and when enabling/disabling in `metrics`.

# Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

### See Also

Other evaluation functions: binomial\_metrics(), confusion\_matrix(), evaluate(), gaussian\_metrics(), multinomial\_metrics()

# **Examples**

```
# Attach packages
library(cvms)

data <- data.frame(
   "targets" = rnorm(100, 14.7, 3.6),
   "predictions" = rnorm(100, 13.2, 4.6)
)

evaluate_residuals(
  data = data,
   target_col = "targets",
   prediction_col = "predictions"
)</pre>
```

56 font

font

Create a list of font settings for plots

# Description

# [Experimental]

Creates a list of font settings for plotting with cvms plotting functions.

NOTE: This is very experimental and will likely change.

# Usage

```
font(
  size = NULL,
 color = NULL,
 alpha = NULL,
  nudge_x = NULL
 nudge_y = NULL,
  angle = NULL,
  family = NULL,
  fontface = NULL,
 hjust = NULL,
  vjust = NULL,
  lineheight = NULL,
 digits = NULL,
 prefix = NULL,
  suffix = NULL
)
```

# **Arguments**

```
size, color, alpha, nudge_x, nudge_y, angle, family, fontface, hjust, vjust, lineheight

As passed to ggplot2::geom_text.

digits

Number of digits to round to. If negative, no rounding will take place.

prefix

A string prefix.

suffix

A string suffix.
```

# Value

List of settings.

## Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

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### See Also

Other plotting functions: plot\_confusion\_matrix(), plot\_metric\_density(), plot\_probabilities(), plot\_probabilities\_ecdf(), sum\_tile\_settings()

gaussian\_metrics

Select metrics for Gaussian evaluation

# Description

### [Experimental]

Enable/disable metrics for Gaussian evaluation. Can be supplied to the `metrics` argument in many of the cvms functions.

Note: Some functions may have slightly different defaults than the ones supplied here.

## Usage

```
gaussian_metrics(
  all = NULL,
  rmse = NULL,
 mae = NULL,
  nrmse_rng = NULL,
  nrmse_iqr = NULL,
  nrmse_std = NULL,
  nrmse_avg = NULL,
  rae = NULL,
  rse = NULL,
  rrse = NULL,
  rmsle = NULL,
 male = NULL,
 mape = NULL,
 mse = NULL,
  tae = NULL,
  tse = NULL,
  r2m = NULL,
  r2c = NULL,
  aic = NULL,
  aicc = NULL,
  bic = NULL
)
```

# **Arguments**

all Enable/disa

Enable/disable all arguments at once. (Logical)

Specifying other metrics will overwrite this, why you can use (all = FALSE, rmse = TRUE) to get only the RMSE metric.

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rmse RMSE. (Default: TRUE) Root Mean Square Error.

mae MAE. (Default: TRUE)

Mean Absolute Error.

nrmse\_rng NRMSE(RNG). (Default: FALSE)

Normalized Root Mean Square Error (by target range).

nrmse\_iqr NRMSE(IQR). (Default: TRUE)

Normalized Root Mean Square Error (by target interquartile range).

nrmse\_std NRMSE(STD). (Default: FALSE)

Normalized Root Mean Square Error (by target standard deviation).

nrmse\_avg NRMSE(AVG). (Default: FALSE)

Normalized Root Mean Square Error (by target mean).

rae RAE. (Default: TRUE)

Relative Absolute Error.

rse RSE. (Default: FALSE)

Relative Squared Error.

rrse RRSE. (Default: TRUE)

Root Relative Squared Error.

rmsle RMSLE. (Default: TRUE)

Root Mean Square Log Error.

male MALE. (Default: FALSE)

Mean Absolute Log Error.

mape MAPE. (Default: FALSE)

Mean Absolute Percentage Error.

mse MSE. (Default: FALSE)

Mean Square Error.

tae TAE. (Default: FALSE)

Total Absolute Error

tse TSE. (Default: FALSE)

Total Squared Error.

r2m r2m. (Default: FALSE)

Marginal R-squared.

r2c r2c. (Default: FALSE)

Conditional R-squared.

aic AIC. (Default: FALSE)

Akaike Information Criterion.

aicc AICc. (Default: FALSE)

Corrected Akaike Information Criterion.

bic BIC. (Default: FALSE)

Bayesian Information Criterion.

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### Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

#### See Also

Other evaluation functions: binomial\_metrics(), confusion\_matrix(), evaluate(), evaluate\_residuals(), multinomial\_metrics()

### **Examples**

```
# Attach packages
library(cvms)

# Enable only RMSE
gaussian_metrics(all = FALSE, rmse = TRUE)

# Enable all but RMSE
gaussian_metrics(all = TRUE, rmse = FALSE)

# Disable RMSE
gaussian_metrics(rmse = FALSE)
```

model\_functions

Examples of model\_fn functions

# Description

# [Experimental]

Examples of model functions that can be used in cross\_validate\_fn(). They can either be used directly or be starting points.

The update\_hyperparameters() function updates the list of hyperparameters with default values for missing hyperparameters. You can also specify required hyperparameters.

### Usage

```
model_functions(name)
```

## **Arguments**

name

Name of model to get model function for, as it appears in the following list:

Name	Function	Hyperparameters (default)
"lm"	stats::lm()	
"lmer"	<pre>lme4::lmer()</pre>	REML (FALSE)
"glm_binomial"	stats::glm()	
"glmer_binomial"	<pre>lme4::glmer()</pre>	

```
"svm_gaussian" e1071::svm() kernel ("radial"), cost (1)

"svm_binomial" e1071::svm() kernel ("radial"), cost (1)

"svm_multinomial" e1071::svm() kernel ("radial"), cost (1)

"naive_bayes" e1071::naiveBayes() laplace (0)
```

#### Value

```
A function with the following form:

function(train_data, formula, hyperparameters) {

# Return fitted model object
}
```

## Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

### See Also

Other example functions: predict\_functions(), preprocess\_functions(), update\_hyperparameters()

most\_challenging

Find the data points that were hardest to predict

# **Description**

[Experimental] Finds the data points that, overall, were the most challenging to predict, based on a prediction metric.

# Usage

```
most_challenging(
  data,
  type,
  obs_id_col = "Observation",
  target_col = "Target",
  prediction_cols = ifelse(type == "gaussian", "Prediction", "Predicted Class"),
  threshold = 0.15,
  threshold_is = "percentage",
  metric = NULL,
  cutoff = 0.5
)
```

# **Arguments**

data

data.frame with predictions, targets and observation IDs. Can be grouped by dplyr::group\_by().

Predictions can be passed as values, predicted classes or predicted probabilities: **N.B.** Adds .Machine\$double.eps to all probabilities to avoid log(0).

**Multinomial:** When `type` is "multinomial", the predictions can be passed in one of two formats.

Probabilities (Preferable):

One column per class with the probability of that class. The columns should have the name of their class, as they are named in the target column. E.g.:

target	class_3	class_2	class_1
class_2	0.203	0.528	0.269
class_3	0.310	0.322	0.368
class_2	0.254	0.371	0.375

#### Classes:

A single column of type character with the predicted classes. E.g.:

prediction	target
class_2	class_2
class_1	class_3
class_1	class_2

**Binomial:** When `type` is "binomial", the predictions can be passed in one of two formats.

*Probabilities (Preferable):* One column with the **probability of class being the second class alphabetically** ("dog" if classes are "cat" and "dog"). E.g.:

prediction	target
0.769	"dog"
0.368	"dog"
0.375	"cat"

Note: At the alphabetical ordering of the class labels, they are of type character, why e.g. 100 would come before 7.

# Classes:

A single column of type character with the predicted classes. E.g.:

prediction	target
class_0	class_1
class_1	class_1
class_1	class_0

**Gaussian:** When `type` is "gaussian", the predictions should be passed as one column with the predicted values. E.g.:

prediction	target
28.9	30.2
33.2	27.1
23.4	21.3

type Type of task used to get the predictions:

"gaussian" for regression (like linear regression).

"binomial" for binary classification.

"multinomial" for multiclass classification.

obs\_id\_col Name of column with observation IDs. This will be used to aggregate the per-

formance of each observation.

target\_col Name of column with the true classes/values in `data`.

prediction\_cols

Name(s) of column(s) with the predictions.

threshold Threshold to filter observations by. Depends on `type` and `threshold\_is`.

The threshold can either be a **percentage** or a **score**. For percentages, a lower threshold returns fewer observations. For scores, this depends on `type`.

### Gaussian:

threshold\_is "percentage": (Approximate) percentage of the observations with the largest root mean square errors to return.

*threshold\_is "score":* Observations with a root mean square error larger than or equal to the threshold will be returned.

### **Binomial, Multinomial:**

threshold\_is "percentage": (Approximate) percentage of the observations to return with:

MAE, Cross Entropy: Highest error scores.

Accuracy: Lowest accuracies

*threshold\_is "score":* MAE, Cross Entropy: Observations with an error score above or equal to the threshold will be returned.

Accuracy: Observations with an accuracy below or equal to the threshold will be returned.

threshold\_is

Either "score" or "percentage". See `threshold`.

metric

The metric to use. If NULL, the default metric depends on the format of the prediction columns.

**Binomial, Multinomial:** "Accuracy", "MAE" or "Cross Entropy".

When *one* prediction column with predicted *classes* is passed, the default is "Accuracy". In this configuration, the other metrics are not calculated.

When *one or more* prediction columns with predicted *probabilities* are passed, the default is "MAE". This is the Mean Absolute Error of the probability of the target class.

Gaussian: Ignored. Always uses "RMSE".

cutoff

Threshold for predicted classes. (Numeric)

N.B. Binomial only.

# Value

data.frame with the most challenging observations and their metrics.

>=` / `<=` denotes the threshold as score.

### Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

# **Examples**

```
# Attach packages
library(cvms)
library(dplyr)
## Multinomial
##
# Find the most challenging data points (per classifier)
# in the predicted.musicians dataset
# which resembles the "Predictions" tibble from the evaluation results
# Passing predicted probabilities
# Observations with 30% highest MAE scores
most_challenging(
  predicted.musicians,
  obs_id_col = "ID",
  prediction_cols = c("A", "B", "C", "D"),
  type = "multinomial",
  threshold = 0.30
)
# Observations with 25% highest Cross Entropy scores
most_challenging(
  predicted.musicians,
  obs_id_col = "ID",
  prediction_cols = c("A", "B", "C", "D"),
  type = "multinomial",
  threshold = 0.25,
  metric = "Cross Entropy"
)
# Passing predicted classes
# Observations with 30% lowest Accuracy scores
most_challenging(
  predicted.musicians,
  obs_id_col = "ID",
  prediction_cols = "Predicted Class",
  type = "multinomial",
  threshold = 0.30
)
```

```
# The 40% lowest-scoring on accuracy per classifier
predicted.musicians %>%
  dplyr::group_by(Classifier) %>%
  most_challenging(
   obs_id_col = "ID",
   prediction_cols = "Predicted Class",
    type = "multinomial",
    threshold = 0.40
  )
# Accuracy scores below 0.05
most_challenging(
  predicted.musicians,
  obs_id_col = "ID",
  type = "multinomial",
  threshold = 0.05,
  threshold_is = "score"
)
##
## Binomial
##
# Subset the predicted.musicians
binom_data <- predicted.musicians %>%
  dplyr::filter(Target %in% c("A","B")) %>%
  dplyr::rename(Prediction = B)
# Passing probabilities
# Observations with 30% highest MAE
most_challenging(
  binom_data,
  obs_id_col = "ID",
  type = "binomial",
  prediction_cols = "Prediction",
  threshold = 0.30
)
# Observations with 30% highest Cross Entropy
most_challenging(
  binom_data,
  obs_id_col = "ID",
  type = "binomial",
  prediction_cols = "Prediction",
  threshold = 0.30,
  metric = "Cross Entropy"
)
# Passing predicted classes
# Observations with 30% lowest Accuracy scores
most_challenging(
  binom_data,
  obs_id_col = "ID",
```

```
type = "binomial",
 prediction_cols = "Predicted Class",
 threshold = 0.30
)
##
## Gaussian
##
set.seed(1)
df <- data.frame(</pre>
 "Observation" = rep(1:10, n = 3),
  "Target" = rnorm(n = 30, mean = 25, sd = 5),
  "Prediction" = rnorm(n = 30, mean = 27, sd = 7)
)
# The 20% highest RMSE scores
most_challenging(
 type = "gaussian",
 threshold = 0.2
)
# RMSE scores above 9
most_challenging(
 type = "gaussian",
 threshold = 9,
 threshold_is = "score"
)
```

multiclass\_probability\_tibble

Generate a multiclass probability tibble

# **Description**

# [Maturing]

Generate a tibble with random numbers containing one column per specified class. When the softmax function is applied, the numbers become probabilities that sum to 1 row-wise. Optionally, add columns with targets and predicted classes.

# Usage

```
multiclass_probability_tibble(
  num_classes,
  num_observations,
```

```
apply_softmax = TRUE,
FUN = runif,
class_name = "class_",
add_predicted_classes = FALSE,
add_targets = FALSE
)
```

### **Arguments**

num\_classes The number of classes. Also the number of columns in the tibble.

num\_observations
The number of observations. Also the number of rows in the tibble.

apply\_softmax
Whether to apply the softmax function row-wise. This will transform the numbers to probabilities that sum to 1 row-wise.

FUN
Function for generating random numbers. The first argument must be the number of random numbers to generate, as no other arguments are supplied.

class\_name
The prefix for the column names. The column index is appended.

add\_predicted\_classes
Whether to add a column with the predicted classes. (Logical)
The class with the highest value is the predicted class.

The class with the highest value is the predicted class.

add\_targets Whether to add a column with randomly selected target classes. (Logical)

# Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

# **Examples**

```
# Attach cvms
library(cvms)
# Create a tibble with 5 classes and 10 observations
# Apply softmax to make sure the probabilities sum to 1
multiclass_probability_tibble(
 num_classes = 5,
 num_observations = 10,
 apply_softmax = TRUE
)
# Using the rnorm function to generate the random numbers
multiclass_probability_tibble(
 num_classes = 5,
 num_observations = 10,
 apply_softmax = TRUE,
 FUN = rnorm
)
# Add targets and predicted classes
multiclass_probability_tibble(
```

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```
num_classes = 5,
 num_observations = 10,
 apply_softmax = TRUE,
 FUN = rnorm,
 add_predicted_classes = TRUE,
 add\_targets = TRUE
)
# Creating a custom generator function that
# exponentiates the numbers to create more "certain" predictions
rcertain <- function(n) {</pre>
  (runif(n, min = 1, max = 100)^1.4) / 100
multiclass_probability_tibble(
 num_classes = 5,
 num_observations = 10,
 apply_softmax = TRUE,
 FUN = rcertain
)
```

multinomial\_metrics

Select metrics for multinomial evaluation

# Description

## [Experimental]

Enable/disable metrics for multinomial evaluation. Can be supplied to the `metrics` argument in many of the cvms functions.

Note: Some functions may have slightly different defaults than the ones supplied here.

# Usage

```
multinomial_metrics(
    all = NULL,
    overall_accuracy = NULL,
    balanced_accuracy = NULL,
    w_balanced_accuracy = NULL,
    accuracy = NULL,
    w_accuracy = NULL,
    f1 = NULL,
    w_f1 = NULL,
    sensitivity = NULL,
    w_sensitivity = NULL,
    specificity = NULL,
    w_specificity = NULL,
    pos_pred_value = NULL,
    w_pos_pred_value = NULL,
```

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```
neg_pred_value = NULL,
     w_neg_pred_value = NULL,
      auc = NULL,
     kappa = NULL,
     w_kappa = NULL,
     mcc = NULL,
     detection_rate = NULL,
     w_detection_rate = NULL,
     detection_prevalence = NULL,
     w_detection_prevalence = NULL,
     prevalence = NULL,
     w_prevalence = NULL,
      false_neg_rate = NULL,
     w_false_neg_rate = NULL,
      false_pos_rate = NULL,
     w_false_pos_rate = NULL,
      false_discovery_rate = NULL,
     w_false_discovery_rate = NULL,
      false_omission_rate = NULL,
     w_false_omission_rate = NULL,
      threat_score = NULL,
     w_threat_score = NULL,
     aic = NULL,
     aicc = NULL,
     bic = NULL
   )
Arguments
   all
                    Enable/disable all arguments at once. (Logical)
                    Specifying other metrics will overwrite this, why you can use (all = FALSE,
                    accuracy = TRUE) to get only the Accuracy metric.
    overall_accuracy
                    Overall Accuracy (Default: TRUE)
    balanced_accuracy
                    Macro Balanced Accuracy (Default: TRUE)
   w_balanced_accuracy
                    Weighted Balanced Accuracy (Default: FALSE)
   accuracy
                    Accuracy (Default: FALSE)
                    Weighted Accuracy (Default: FALSE)
   w_accuracy
    f1
                    F1 (Default: TRUE)
   w_f1
                    Weighted F1 (Default: FALSE)
                    Sensitivity (Default: TRUE)
    sensitivity
                    Weighted Sensitivity (Default: FALSE)
   w_sensitivity
                    Specificity (Default: TRUE)
    specificity
                    Weighted Specificity (Default: FALSE)
   w_specificity
```

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pos\_pred\_value Pos Pred Value (Default: TRUE) w\_pos\_pred\_value Weighted Pos Pred Value (Default: FALSE) neg\_pred\_value Neg Pred Value (Default: TRUE) w\_neg\_pred\_value Weighted Neg Pred Value (Default: FALSE) auc AUC (Default: FALSE) Kappa (Default: TRUE) kappa Weighted Kappa (Default: FALSE) w\_kappa MCC (Default: TRUE) mcc Multiclass Matthews Correlation Coefficient. detection\_rate Detection Rate (Default: TRUE) w\_detection\_rate Weighted Detection Rate (Default: FALSE) detection\_prevalence Detection Prevalence (Default: TRUE) w\_detection\_prevalence Weighted Detection Prevalence (Default: FALSE) Prevalence (Default: TRUE) prevalence w\_prevalence Weighted Prevalence (Default: FALSE) false\_neg\_rate False Neg Rate (Default: FALSE) w\_false\_neg\_rate Weighted False Neg Rate (Default: FALSE) false\_pos\_rate False Pos Rate (Default: FALSE) w\_false\_pos\_rate Weighted False Pos Rate (Default: FALSE) false\_discovery\_rate False Discovery Rate (Default: FALSE) w\_false\_discovery\_rate Weighted False Discovery Rate (Default: FALSE) false\_omission\_rate False Omission Rate (Default: FALSE) w\_false\_omission\_rate Weighted False Omission Rate (Default: FALSE) threat\_score Threat Score (Default: FALSE) w\_threat\_score Weighted Threat Score (Default: FALSE) aic AIC. (Default: FALSE) AICc. (Default: FALSE) aicc bic BIC. (Default: FALSE)

## Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

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### See Also

Other evaluation functions: binomial\_metrics(), confusion\_matrix(), evaluate(), evaluate\_residuals(), gaussian\_metrics()

### **Examples**

```
# Attach packages
library(cvms)

# Enable only Balanced Accuracy
multinomial_metrics(all = FALSE, balanced_accuracy = TRUE)

# Enable all but Balanced Accuracy
multinomial_metrics(all = TRUE, balanced_accuracy = FALSE)

# Disable Balanced Accuracy
multinomial_metrics(balanced_accuracy = FALSE)
```

musicians

Musician groups

### **Description**

Made-up data on 60 musicians in 4 groups for multiclass classification.

### Format

A data.frame with 60 rows and 9 variables:

**ID** Musician identifier, 60 levels

Age Age of the musician. Between 17 and 66 years.

Class The class of the musician. One of "A", "B", "C", and "D".

Height Height of the musician. Between 146 and 196 centimeters.

**Drums** Whether the musician plays drums. 0 = No, 1 = Yes.

**Bass** Whether the musician plays bass. 0 = No, 1 = Yes.

**Guitar** Whether the musician plays guitar. 0 = No, 1 = Yes.

**Keys** Whether the musician plays keys. 0 = No, 1 = Yes.

**Vocals** Whether the musician sings. 0 = No, 1 = Yes.

#### Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

### See Also

predicted.musicians

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participant.scores

Participant scores

### **Description**

Made-up experiment data with 10 participants and two diagnoses. Test scores for 3 sessions per participant, where participants improve their scores each session.

### **Format**

A data. frame with 30 rows and 5 variables:

participant participant identifier, 10 levels

age age of the participant, in years

diagnosis diagnosis of the participant, either 1 or 0

**score** test score of the participant, on a 0-100 scale

session testing session identifier, 1 to 3

### Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

plot\_confusion\_matrix Plot a confusion matrix

#### **Description**

#### [Experimental]

Creates a ggplot2 object representing a confusion matrix with counts, overall percentages, row percentages and column percentages. An extra row and column with sum tiles and the total count can be added.

The confusion matrix can be created with evaluate(). See `Examples`.

While this function is intended to be very flexible (hence the large number of arguments), the defaults should work in most cases for most users. See the Examples.

**NEW**: Our **Plot Confusion Matrix** web application allows using this function without code. Select from multiple design templates or make your own.

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# Usage

```
plot_confusion_matrix(
  conf_matrix,
  target_col = "Target",
 prediction_col = "Prediction",
  counts_col = "N",
  sub\_col = NULL,
  class_order = NULL,
  add_sums = FALSE,
  add_counts = TRUE,
  add_normalized = TRUE,
  add_row_percentages = TRUE,
  add_col_percentages = TRUE,
  diag_percentages_only = FALSE,
  rm_zero_percentages = TRUE,
  rm_zero_text = TRUE,
  add_zero_shading = TRUE,
  amount_3d_effect = 1,
  add_arrows = TRUE,
  counts_on_top = FALSE,
  palette = "Blues",
  intensity_by = "counts",
  intensity_lims = NULL,
  intensity_beyond_lims = "truncate",
  theme_fn = ggplot2::theme_minimal,
  place_x_axis_above = TRUE,
  rotate_y_text = TRUE,
  digits = 1,
  font_counts = font(),
  font_normalized = font(),
  font_row_percentages = font(),
  font_col_percentages = font(),
  arrow_size = 0.048,
  arrow_nudge_from_text = 0.065,
  tile_border_color = NA,
  tile_border_size = 0.1,
  tile_border_linetype = "solid",
  sums_settings = sum_tile_settings(),
  darkness = 0.8
)
```

## **Arguments**

 $conf\_matrix$ 

Confusion matrix tibble with each combination of targets and predictions along with their counts.

E.g. for a binary classification:

# Target Prediction N

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class_1	class_1	5
class_1	class_2	9
class_2	class_1	3
class 2	class 2	2

As created with the various evaluation functions in cvms, like evaluate().

An additional `sub\_col` column (character) can be specified as well. Its content will replace the bottom text ('counts' by default or 'normalized' when `counts\_on\_top` is enabled).

**Note**: If you supply the results from evaluate() or confusion\_matrix() directly, the confusion matrix tibble is extracted automatically, if possible.

target\_col Name of column with target levels.

prediction\_col Name of column with prediction levels.

counts\_col Name of column with a count for each combination of the target and prediction

levels.

sub\_col Name of column with text to replace the bottom text ('counts' by default or

'normalized' when `counts\_on\_top` is enabled).

It simply replaces the text, so all settings will still be called e.g. `font\_counts` etc. When other settings make it so, that no bottom text is displayed (e.g.  $\frac{1}{2}$ ).

`add\_counts` = FALSE), this text is not displayed either.

class\_order Names of the classes in `conf\_matrix` in the desired order. When NULL, the

classes are ordered alphabetically. Note that the entire set of unique classes from

both `target\_col` and `prediction\_col` must be specified.

add\_sums Add tiles with the row/column sums. Also adds a total count tile. (Logical)

The appearance of these tiles can be specified in `sums\_settings`.

Note: Adding the sum tiles with a palette requires the ggnewscale package.

add\_counts Add the counts to the middle of the tiles. (Logical)

add\_normalized Normalize the counts to percentages and add to the middle of the tiles. (Logical) add\_row\_percentages

Add the row percentages, i.e. how big a part of its row the tile makes up. (Logical)

By default, the row percentage is placed to the right of the tile, rotated 90 degrees.

add\_col\_percentages

Add the column percentages, i.e. how big a part of its column the tile makes up. (Logical)

By default, the row percentage is placed at the bottom of the tile.

diag\_percentages\_only

Whether to only have row and column percentages in the diagonal tiles. (Logical)

rm\_zero\_percentages

Whether to remove row and column percentages when the count is 0. (Logical)

digits

no rounding.

Whether to remove counts and normalized percentages when the count is 0. rm\_zero\_text (Logical) add\_zero\_shading Add image of skewed lines to zero-tiles. (Logical) Note: Adding the zero-shading requires the rsvg and ggimage packages. amount\_3d\_effect Amount of 3D effect (tile overlay) to add. Passed as whole number from 0 (no effect) up to 6 (biggest effect). This helps separate tiles with the same intensities. Note: The overlay may not fit the tiles in many-class cases that haven't been tested. If the boxes do not overlap properly, simply turn it off. add\_arrows Add the arrows to the row and col percentages. (Logical) Note: Adding the arrows requires the rsvg and ggimage packages. Switch the counts and normalized counts, such that the counts are on top. (Logcounts\_on\_top ical) Color scheme. Passed directly to `palette` in ggplot2::scale\_fill\_distiller. palette Try these palettes: "Greens", "Oranges", "Greys", "Purples", "Reds", as well as the default "Blues". Alternatively, pass a named list with limits of a custom gradient as e.g. `list("low"="#B1F9E8", "high"="#239895")`. These are passed to ggplot2::scale\_fill\_gradient. intensity\_by The measure that should control the color intensity of the tiles. Either `counts`, `normalized` or one of `log counts`, `log2 counts`, `log10 counts`, `arcsinh counts`. For 'normalized', the color limits become 0-100 (except when `intensity\_lims` are specified), why the intensities can better be compared across plots. For the 'log\*' and 'arcsinh' versions, the log/arcsinh transformed counts are used. Note: In 'log\*' transformed counts, 0-counts are set to '0', why they won't be distinguishable from 1-counts. intensity\_lims A specific range of values for the color intensity of the tiles. Given as a numeric vector with c(min, max). This allows having the same intensity scale across plots for better comparison of prediction sets. intensity\_beyond\_lims What to do with values beyond the `intensity\_lims`. One of "truncate", "grey". theme\_fn The ggplot2 theme function to apply. place\_x\_axis\_above Move the x-axis text to the top and reverse the levels such that the "correct" diagonal goes from top left to bottom right. (Logical) Whether to rotate the y-axis text to be vertical instead of horizontal. (Logical) rotate\_y\_text

Number of digits to round to (percentages only). Set to a negative number for

Can be set for each font individually via the font\_\* arguments.

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```
font_counts
                 list of font settings for the counts. Can be provided with font().
font_normalized
                 list of font settings for the normalized counts. Can be provided with font().
font_row_percentages
                 list of font settings for the row percentages. Can be provided with font().
font_col_percentages
                 list of font settings for the column percentages. Can be provided with font().
arrow_size
                 Size of arrow icons. (Numeric)
                 Is divided by sqrt(nrow(conf_matrix)) and passed on to ggimage::geom_icon().
arrow_nudge_from_text
                 Distance from the percentage text to the arrow. (Numeric)
tile_border_color
                 Color of the tile borders. Passed as 'colour' to ggplot2::geom_tile.
tile_border_size
                  Size of the tile borders. Passed as 'size' to ggplot2::geom_tile.
tile_border_linetype
                 Linetype for the tile borders. Passed as 'linetype' to ggplot2::geom_tile.
sums_settings
                 A list of settings for the appearance of the sum tiles. Can be provided with
                 sum_tile_settings().
darkness
                 How dark the darkest colors should be, between 0 and 1, where 1 is darkest.
                 Technically, a lower value increases the upper limit in ggplot2::scale_fill_distiller.
```

#### **Details**

Inspired by Antoine Sachet's answer at https://stackoverflow.com/a/53612391/11832955

#### Value

A ggplot2 object representing a confusion matrix. Color intensity depends on either the counts (default) or the overall percentages.

By default, each tile has the normalized count (overall percentage) and count in the middle, the column percentage at the bottom, and the row percentage to the right and rotated 90 degrees.

In the "correct" diagonal (upper left to bottom right, by default), the column percentages are the class-level sensitivity scores, while the row percentages are the class-level positive predictive values.

# Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

#### See Also

```
Other plotting functions: font(), plot_metric_density(), plot_probabilities(), plot_probabilities_ecdf(), sum_tile_settings()
```

# **Examples**

```
# Attach cvms
library(cvms)
library(ggplot2)
# Two classes
# Create targets and predictions data frame
data <- data.frame(</pre>
 "B", "B", "A", "B", "A", "A", "A", "A"),
 stringsAsFactors = FALSE
)
# Evaluate predictions and create confusion matrix
evaluation <- evaluate(</pre>
 data = data,
 target_col = "target",
 prediction_cols = "prediction",
 type = "binomial"
)
# Inspect confusion matrix tibble
evaluation[["Confusion Matrix"]][[1]]
# Plot confusion matrix
# Supply confusion matrix tibble directly
plot_confusion_matrix(evaluation[["Confusion Matrix"]][[1]])
# Plot first confusion matrix in evaluate() output
plot_confusion_matrix(evaluation)
## Not run:
# Add sum tiles
plot_confusion_matrix(evaluation, add_sums = TRUE)
## End(Not run)
# Add labels to diagonal row and column percentages
# This example assumes "B" is the positive class
# but you could write anything as prefix to the percentages
plot_confusion_matrix(
   evaluation,
    font_row_percentages = font(prefix=c("NPV = ", "", "", "PPV = ")),
    font_col_percentages = font(prefix=c("Spec = ", "", "", "Sens = "))
)
# Three (or more) classes
```

plot\_confusion\_matrix

```
# Create targets and predictions data frame
data <- data.frame(</pre>
  "target" = c("A", "B", "C", "B", "A", "B", "C",

"B", "A", "B", "C", "B", "A"),

"prediction" = c("C", "B", "A", "C", "A", "B", "B",

"C", "A", "B", "C", "A", "C"),
  stringsAsFactors = FALSE
)
# Evaluate predictions and create confusion matrix
evaluation <- evaluate(</pre>
  data = data,
  target_col = "target",
  prediction_cols = "prediction",
  type = "multinomial"
# Inspect confusion matrix tibble
evaluation[["Confusion Matrix"]][[1]]
# Plot confusion matrix
# Supply confusion matrix tibble directly
plot_confusion_matrix(evaluation[["Confusion Matrix"]][[1]])
# Plot first confusion matrix in evaluate() output
plot_confusion_matrix(evaluation)
## Not run:
# Add sum tiles
plot_confusion_matrix(evaluation, add_sums = TRUE)
## End(Not run)
# Counts only
plot_confusion_matrix(
  evaluation[["Confusion Matrix"]][[1]],
  add_normalized = FALSE,
  add_row_percentages = FALSE,
  add_col_percentages = FALSE
)
# Change color palette to green
# Change theme to `theme_light`.
plot_confusion_matrix(
  evaluation[["Confusion Matrix"]][[1]],
  palette = "Greens",
  theme_fn = ggplot2::theme_light
)
## Not run:
# Change colors palette to custom gradient
```

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```
# with a different gradient for sum tiles
plot_confusion_matrix(
  evaluation[["Confusion Matrix"]][[1]],
  palette = list("low" = "#B1F9E8", "high" = "#239895"),
  sums_settings = sum_tile_settings(
    palette = list("low" = "#e9e1fc", "high" = "#BE94E6")
  add\_sums = TRUE
)
## End(Not run)
# The output is a ggplot2 object
# that you can add layers to
# Here we change the axis labels
plot_confusion_matrix(evaluation[["Confusion Matrix"]][[1]]) +
  ggplot2::labs(x = "True", y = "Guess")
# Replace the bottom tile text
# with some information
# First extract confusion matrix
# Then add new column with text
cm <- evaluation[["Confusion Matrix"]][[1]]</pre>
cm[["Trials"]] <- c(</pre>
  "(8/9)", "(3/9)", "(1/9)",
"(3/9)", "(7/9)", "(4/9)",
"(1/9)", "(2/9)", "(8/9)"
# Now plot with the `sub_col` argument specified
plot_confusion_matrix(cm, sub_col="Trials")
```

plot\_metric\_density

Density plot for a metric

## **Description**

#### [Experimental]

Creates a ggplot2 object with a density plot for one of the columns in the passed data.frame(s).

Note: In its current form, it is mainly intended as a quick way to visualize the results from cross-validations and baselines (random evaluations). It may change significantly in future versions.

#### Usage

```
plot_metric_density(
  results = NULL,
```

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```
baseline = NULL,
metric = "",
fill = c("darkblue", "lightblue"),
alpha = 0.6,
theme_fn = ggplot2::theme_minimal,
xlim = NULL
)
```

# **Arguments**

results data. frame with a metric column to create density plot for.

To only plot the baseline, set to NULL.

baseline data.frame with the random evaluations from baseline(). Should contain a

column for the metric.

To only plot the results, set to NULL.

metric Name of the metric column in `results` to plot. (Character)

fill Colors of the plotted distributions. The first color is for the `baseline`, the

second for the `results`.

alpha Transparency of the distribution (0 - 1). theme\_fn The ggplot2 theme function to apply. xlim Limits for the x-axis. Can be set to NULL.

E.g. c(0, 1).

# Value

A ggplot2 object with the density of a metric, possibly split in 'Results' and 'Baseline'.

# Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

#### See Also

```
Other plotting functions: font(), plot_confusion_matrix(), plot_probabilities(), plot_probabilities_ecdf(), sum_tile_settings()
```

# Examples

```
# Attach packages
library(cvms)
library(dplyr)

# We will use the musicians and predicted.musicians datasets
musicians
predicted.musicians

# Set seed
set.seed(42)
```

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```
# Create baseline for targets
bsl <- baseline_multinomial(</pre>
  test_data = musicians,
  dependent_col = "Class",
  n = 20 # Normally 100
)
# Evaluate predictions grouped by classifier and fold column
eval <- predicted.musicians %>%
  dplyr::group_by(Classifier, `Fold Column`) %>%
  evaluate(
  target_col = "Target",
  prediction_cols = c("A", "B", "C", "D"),
  type = "multinomial"
# Plot density of the Overall Accuracy metric
plot_metric_density(
  results = eval,
  baseline = bsl$random_evaluations,
  metric = "Overall Accuracy",
  xlim = c(0,1)
)
# The bulk of classifier results are much better than
# the baseline results
```

precomputed.formulas Precomputed formulas

# **Description**

Fixed effect combinations for model formulas with/without two- and three-way interactions. Up to eight fixed effects in total with up to five fixed effects per formula.

#### **Format**

```
A data.frame with 259,358 rows and 5 variables:
```

formula\_ combination of fixed effects, separated by "+" and "\*"

max\_interaction\_size maximum interaction size in the formula, up to 3

**max\_effect\_frequency** maximum count of an effect in the formula, e.g. the 3 A's in "A  $\star$  B + A  $\star$  C + A  $\star$  D"

**num\_effects** number of unique effects included in the formula

min\_num\_fixed\_effects minimum number of fixed effects required to use the formula, i.e. the index in the alphabet of the last of the alphabetically ordered effects (letters) in the formula, so 4 for the formula: "A + B + D"

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## **Details**

Effects are represented by the first eight capital letters.

Used by combine\_predictors.

#### Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

predicted.musicians Predicted musician groups

# **Description**

Predictions by 3 classifiers of the 4 classes in the musicians dataset. Obtained with 5-fold stratified cross-validation (3 repetitions). The three classifiers were fit using nnet::multinom, randomForest::randomForest, and e1071::svm.

#### **Format**

A data. frame with 540 rows and 10 variables:

Classifier The applied classifier. One of "nnet\_multinom", "randomForest", and "e1071\_svm".

**Fold Column** The fold column name. Each is a unique 5-fold split. One of ".folds\_1", ".folds\_2", and ".folds\_3".

**Fold** The fold. 1 to 5.

**ID** Musician identifier, 60 levels

**Target** The actual class of the musician. One of "A", "B", "C", and "D".

- **A** The probability of class "A".
- **B** The probability of class "B".
- **C** The probability of class "C".
- **D** The probability of class "D".

**Predicted Class** The predicted class. The argmax of the four probability columns.

#### **Details**

```
Used formula: "Class ~ Height + Age + Drums + Bass + Guitar + Keys + Vocals"
```

## Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

## See Also

musicians

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## **Examples**

```
# Attach packages
library(cvms)
library(dplyr)
# Evaluate each fold column
predicted.musicians %>%
  dplyr::group_by(Classifier, `Fold Column`) %>%
  evaluate(target_col = "Target",
           prediction_cols = c("A", "B", "C", "D"),
           type = "multinomial")
# Overall ID evaluation
# I.e. if we average all 9 sets of predictions,
# how well did we predict the targets?
overall_id_eval <- predicted.musicians %>%
  evaluate(target_col = "Target",
           prediction_cols = c("A", "B", "C", "D"),
           type = "multinomial",
           id_col = "ID")
overall_id_eval
# Plot the confusion matrix
plot_confusion_matrix(overall_id_eval$`Confusion Matrix`[[1]])
```

predict\_functions

Examples of predict\_fn functions

# **Description**

## [Experimental]

Examples of predict functions that can be used in cross\_validate\_fn(). They can either be used directly or be starting points.

## Usage

```
predict_functions(name)
```

## **Arguments**

name

Name of model to get predict function for, as it appears in the following table.

The **Model HParams** column lists hyperparameters used in the respective model function.

Name	Function
"lm"	stats::lm()
"lmer"	<pre>lme4::lmer()</pre>

**Model HParams** 

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```
"glm_binomial"
                                            stats::glm()
                                                                                     family = "binomial"
         "glmer_binomial"
                                           lme4::glmer()
                                                                                     family = "binomial"
           "svm gaussian"
                                            e1071::svm()
                                                                                 type = "eps-regression"
           "svm_binomial"
                                                           type = "C-classification", probability = TRUE
                                            e1071::svm()
        "svm multinomial"
                                            e1071::svm()
                                                           type = "C-classification", probability = TRUE
            "naive_bayes"
                                    e1071::naiveBayes()
          "nnet multinom"
                                        nnet::multinom()
           "nnet gaussian"
                                                                                           linout = TRUE
                                            nnet::nnet()
          "nnet binomial"
                                            nnet::nnet()
  "randomForest_gaussian"
                           randomForest::randomForest()
  "randomForest binomial"
                           randomForest::randomForest()
"randomForest_multinomial"
                           randomForest()
```

#### Value

```
A function with the following form:
function(test_data, model, formula, hyperparameters, train_data) {
    # Use model to predict test_data
    # Return predictions
}
```

#### Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

#### See Also

Other example functions: model\_functions(), preprocess\_functions(), update\_hyperparameters()

# **Description**

# [Experimental]

Examples of preprocess functions that can be used in cross\_validate\_fn() and validate\_fn(). They can either be used directly or be starting points.

The examples use recipes, but you can also use caret::preProcess() or similar functions.

In these examples, the preprocessing will only affect the numeric predictors.

You may prefer to hardcode a formula like " $y \sim .$ " (where y is your dependent variable) as that will allow you to set 'preprocess\_one' to TRUE in cross\_validate\_fn() and validate\_fn() and save time.

# Usage

```
preprocess_functions(name)
```

## **Arguments**

name

Name of preprocessing function as it appears in the following list:

Name	Description
"standardize"	Centers and scales the numeric predictors
"range"	Normalizes the numeric predictors to the 0-1 range
"scale"	Scales the numeric predictors to have a standard deviation of one
"center"	Centers the numeric predictors to have a mean of zero
"warn"	Identity function that throws a warning and a message

#### Value

```
A function with the following form:
```

```
function(train_data, test_data, formula, hyperparameters) {
    # Preprocess train_data and test_data
    # Return a list with the preprocessed datasets
    # and optionally a data frame with preprocessing parameters
    list(
        "train" = train_data,
        "test" = test_data,
        "parameters" = tidy_parameters
    )
}
```

# Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

# See Also

Other example functions: model\_functions(), predict\_functions(), update\_hyperparameters()

process\_info\_binomial A set of process information object constructors

# **Description**

# [Experimental]

Classes for storing process information from prediction evaluations.

Used internally.

# Usage

```
process_info_binomial(
  data,
  target_col,
  prediction_cols,
  id_col,
  cat_levels,
 positive,
  cutoff,
  locale = NULL
)
## S3 method for class 'process_info_binomial'
print(x, ...)
## S3 method for class 'process_info_binomial'
as.character(x, ...)
process_info_multinomial(
  data,
  target_col,
  prediction_cols,
 pred_class_col,
  id_col,
  cat_levels,
  apply_softmax,
  locale = NULL
## S3 method for class 'process_info_multinomial'
print(x, ...)
## S3 method for class 'process_info_multinomial'
as.character(x, ...)
process_info_gaussian(data, target_col, prediction_cols, id_col, locale = NULL)
## S3 method for class 'process_info_gaussian'
print(x, ...)
## S3 method for class 'process_info_gaussian'
as.character(x, ...)
```

# **Arguments**

data Data frame.
target\_col Name of target column.

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prediction\_cols

Names of prediction columns.

id\_col Name of ID column.

cat\_levels Categorical levels (classes).
positive Name of the positive class.

cutoff The cutoff used to get class predictions from probabilities.

locale The locale when performing the evaluation. Relevant when any sorting has been

performed.

x a process info object used to select a method.

. . . further arguments passed to or from other methods.

pred\_class\_col Name of predicted classes column.
apply\_softmax Whether softmax has been applied.

#### Value

List with relevant information.

# Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

## **Description**

## [Maturing]

In the (cross-)validation results from functions like cross\_validate(), the model formulas have been split into the columns Dependent, Fixed and Random. Quickly reconstruct the model formulas from these columns.

# Usage

```
reconstruct_formulas(results, topn = NULL)
```

## **Arguments**

results data.frame with results from cross\_validate() or validate(). (tbl)

Must contain at least the columns "Dependent" and "Fixed". For random ef-

fects, the "Random" column should be included.

topn Number of top rows to return. Simply applies head() to the results tibble.

## Value

list of model formulas.

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#### Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

select\_definitions

Select model definition columns

#### **Description**

#### [Experimental]

Select the columns that define the models, such as the formula terms and hyperparameters.

If an expected column is not in the `results` tibble, it is simply ignored.

#### Usage

```
select_definitions(results, unnest_hparams = TRUE, additional_includes = NULL)
```

# Arguments

```
results Results tibble. E.g. from cross_validate() or evaluate().
unnest_hparams Whether to unnest the HParams column. (Logical)
additional_includes
Names of additional columns to select. (Character)
```

#### Value

The model definition columns from the results tibble.

#### Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

select\_metrics

Select columns with evaluation metrics and model definitions

## **Description**

## [Maturing]

When reporting results, we might not want all the nested tibbles and process information columns. This function selects the evaluation metrics and model formulas only.

If an expected column is not in the `results` tibble, it is simply ignored.

## Usage

```
select_metrics(results, include_definitions = TRUE, additional_includes = NULL)
```

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# **Arguments**

```
results Results tibble. E.g. from cross_validate() or evaluate().

include_definitions

Whether to include the Dependent, Fixed and (possibly) Random and HParams columns. (Logical)

additional_includes

Names of additional columns to select. (Character)
```

#### Value

The results tibble with only the metric and model definition columns.

#### Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

simplify\_formula

Simplify formula with inline functions

# **Description**

## [Experimental]

Extracts all variables from a formula object and creates a new formula with all predictor variables added together without the inline functions.

```
E.g.:

y \sim x*z + \log(a) + (1|b)

becomes

y \sim x + z + a + b.
```

This is useful when passing a formula to recipes::recipe() for preprocessing a dataset, as used in the preprocess\_functions().

## Usage

```
simplify_formula(formula, data = NULL, string_out = FALSE)
```

# **Arguments**

formula Formula object.

If a string is passed, it will be converted with as.formula().

When a side *only* contains a NULL, it is kept. Otherwise NULLs are removed. An intercept (1) will only be kept if there are no variables on that side of the

formula.

data data. frame. Used to extract variables when the formula contains a ".".

string\_out Whether to return as a string. (Logical)

summarize\_metrics 89

## Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

## **Examples**

```
# Attach cvms
library(cvms)

# Create formula
f1 <- "y ~ x*z + log(a) + (1|b)"

# Simplify formula (as string)
simplify_formula(f1)

# Simplify formula (as formula)
simplify_formula(as.formula(f1))</pre>
```

summarize\_metrics

Summarize metrics with common descriptors

# **Description**

## [Experimental]

Summarizes all numeric columns. Counts the NAs and Infs in the columns.

# Usage

```
summarize_metrics(data, cols = NULL, na.rm = TRUE, inf.rm = TRUE)
```

#### **Arguments**

data	data. frame with numeric columns to summarize.
cols	Names of columns to summarize. Non-numeric columns are ignored. (Character)
na.rm	Whether to remove NAs before summarizing. (Logical)
inf.rm	Whether to remove Infs before summarizing. (Logical)

## Value

tibble where each row is a descriptor of the column.

The **Measure** column contains the name of the descriptor.

The **NAs** row is a count of the NAs in the column.

The **INFs** row is a count of the Infs in the column.

# Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

90 sum\_tile\_settings

# **Examples**

sum\_tile\_settings

Create a list of settings for the sum tiles in plot\_confusion\_matrix()

# Description

# [Experimental]

Creates a list of settings for plotting the column/row sums in plot\_confusion\_matrix().

The `tc\_` in the arguments refers to the **total count** tile.

NOTE: This is very experimental and will likely change.

# Usage

```
sum_tile_settings(
 palette = NULL,
 label = NULL,
  tile_fill = NULL,
  font_color = NULL,
  tile_border_color = NULL,
  tile_border_size = NULL,
  tile_border_linetype = NULL,
  tc_tile_fill = NULL,
  tc_font_color = NULL,
  tc_tile_border_color = NULL,
  tc_tile_border_size = NULL,
  tc_tile_border_linetype = NULL,
  intensity_by = NULL,
  intensity_lims = NULL,
  intensity_beyond_lims = NULL
)
```

91 sum\_tile\_settings

#### **Arguments**

palette Color scheme to use for sum tiles. Should be different from the `palette` used for the regular tiles. Passed directly to `palette` in ggplot2::scale\_fill\_distiller. Try these palettes: "Greens", "Oranges", "Greys", "Purples", "Reds", and "Blues". Alternatively, pass a named list with limits of a custom gradient as e.g. `list("low"="#e9e1fc", "high"="#BE94E6")`. These are passed to ggplot2::scale\_fill\_gradient. Note: When `tile\_fill` is specified, the `palette` is **ignored**. label The label to use for the sum column and the sum row. tc\_tile\_fill, tile\_fill Specific background color for the tiles. Passed as 'fill' to ggplot2::geom\_tile. If specified, the `palette` is ignored. tc\_font\_color, font\_color Color of the text in the tiles with the column and row sums. tc\_tile\_border\_color, tile\_border\_color Color of the tile borders. Passed as 'colour' to ggplot2::geom\_tile. tc\_tile\_border\_size, tile\_border\_size Size of the tile borders. Passed as 'size' to ggplot2::geom\_tile. tc\_tile\_border\_linetype, tile\_border\_linetype Linetype for the tile borders. Passed as 'linetype' to ggplot2::geom\_tile. The measure that should control the color intensity of the tiles. Either `counts`, intensity\_by `normalized` or one of `log counts`, `log2 counts`, `log10 counts`, `arcsinh counts`. For 'normalized', the color limits become 0-100 (except when `intensity\_lims` are specified), why the intensities can better be compared across plots. For the 'log\*' and 'arcsinh' versions, the log/arcsinh transformed counts are used. **Note**: In 'log\*' transformed counts, 0-counts are set to '0', why they won't be distinguishable from 1-counts. intensity\_lims A specific range of values for the color intensity of the tiles. Given as a numeric vector with c(min, max).

> This allows having the same intensity scale across plots for better comparison of prediction sets.

intensity\_beyond\_lims

What to do with values beyond the `intensity\_lims`. One of "truncate", "grev".

## Value

List of settings.

## Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

## See Also

Other plotting functions: font(), plot\_confusion\_matrix(), plot\_metric\_density(), plot\_probabilities(), plot\_probabilities\_ecdf()

update\_hyperparameters

Check and update hyperparameters

# **Description**

## [Experimental]

- 1. Checks if the required hyperparameters are present and throws an error when it is not the case.
- 2. Inserts the missing hyperparameters with the supplied default values.

For managing hyperparameters in custom model functions for cross\_validate\_fn() or validate\_fn().

## Usage

```
update_hyperparameters(..., hyperparameters, .required = NULL)
```

#### **Arguments**

... Default values for missing hyperparameters.

E.g.:

kernel = "linear", cost = 10

hyperparameters

list of hyperparameters as supplied to  ${\tt cross\_validate\_fn()}$ . Can also be a

single-row data. frame.

required Names of required hyperparameters. If any of these are not present in the hy-

perparameters, an error is thrown.

# Value

A named list with the updated hyperparameters.

## Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

#### See Also

Other example functions: model\_functions(), predict\_functions(), preprocess\_functions()

## **Examples**

```
# Attach packages
library(cvms)
# Create a list of hyperparameters
hparams <- list(</pre>
  "kernel" = "radial",
  "scale" = TRUE
)
# Update hyperparameters with defaults
# Only 'cost' is changed as it's missing
update_hyperparameters(
  cost = 10,
  kernel = "linear",
  "scale" = FALSE,
  hyperparameters = hparams
)
# 'cost' is required
# throws error
if (requireNamespace("xpectr", quietly = TRUE)){
  xpectr::capture_side_effects(
   update_hyperparameters(
      kernel = "linear",
      "scale" = FALSE,
      hyperparameters = hparams,
      .required = "cost"
 )
}
```

validate

Validate regression models on a test set

# Description

# [Stable]

Train linear or logistic regression models on a training set and validate it by predicting a test/validation set. Returns results in a tibble for easy reporting, along with the trained models.

See validate\_fn() for use with custom model functions.

#### Usage

```
validate(
  train_data,
```

```
formulas,
  family,
  test_data = NULL,
  partitions_col = ".partitions",
  control = NULL,
 REML = FALSE,
  cutoff = 0.5,
  positive = 2,
 metrics = list(),
  preprocessing = NULL,
  err_nc = FALSE,
  rm_nc = FALSE,
  parallel = FALSE,
  verbose = FALSE,
  link = deprecated(),
 models = deprecated(),
 model_verbose = deprecated()
)
```

# **Arguments**

train\_data data.frame.

Can contain a grouping factor for identifying partitions - as made with groupdata2::partition().

See `partitions\_col`.

formulas Model formulas as strings. (Character)

E.g. c("y~x", "y~z"). Can contain random effects.

E.g.  $c("y\sim x+(1|r)", "y\sim z+(1|r)")$ .

family Name of the family. (Character)

Currently supports "gaussian" for linear regression with lm() / lme4::lmer()

and "binomial" for binary classification with glm() / lme4::glmer().

See cross\_validate\_fn() for use with other model functions.

test\_data data.frame. If specifying `partitions\_col`, this can be NULL.

partitions\_col Name of grouping factor for identifying partitions. (Character)

Rows with the value 1 in `partitions\_col` are used as training set and rows

with the value 2 are used as test set.

N.B. Only used if 'test\_data' is NULL.

control Construct control structures for mixed model fitting (with lme4::lmer() or

lme4::glmer()). See lme4::lmerControl and lme4::glmerControl.

N.B. Ignored if fitting lm() or glm() models.

REML Restricted Maximum Likelihood. (Logical)

cutoff Threshold for predicted classes. (Numeric)

N.B. Binomial models only

positive

Level from dependent variable to predict. Either as character (*preferable*) or level index (1 or 2 - alphabetically).

E.g. if we have the levels "cat" and "dog" and we want "dog" to be the positive class, we can either provide "dog" or 2, as alphabetically, "dog" comes after "cat".

**Note:** For *reproducibility*, it's preferable to **specify the name directly**, as different locales may sort the levels differently.

Used when calculating confusion matrix metrics and creating ROC curves.

The Process column in the output can be used to verify this setting.

N.B. Only affects evaluation metrics, not the model training or returned predictions.

#### N.B. Binomial models only.

metrics

list for enabling/disabling metrics.

E.g. list("RMSE" = FALSE) would remove RMSE from the results, and list("Accuracy" = TRUE) would add the regular Accuracy metric to the classification results. De-

fault values (TRUE/FALSE) will be used for the remaining available metrics.

You can enable/disable all metrics at once by including "all" = TRUE/FALSE in the list. This is done prior to enabling/disabling individual metrics, why list("all" = FALSE, "RMSE" = TRUE) would return only the RMSE metric.

The list can be created with gaussian\_metrics() or binomial\_metrics().

Also accepts the string "all".

preprocessing

Name of preprocessing to apply.

Available preprocessings are:

Name

"standardize"

Centers and scales the numeri

"range" "scale"

Normalizes the numeric predictors to the 0-1 range. Values outside the min/max range in the test fold are trun

Scales the numeric predictors to have a standard deviation Centers the numeric predictors to have a m

"center"

The preprocessing parameters (mean, SD, etc.) are extracted from the training folds and applied to both the training folds and the test fold. They are returned in the **Preprocess** column for inspection.

N.B. The preprocessings should not affect the results to a noticeable degree, although "range" might due to the truncation.

Whether to message process information like the number of model instances to

err\_nc

Whether to raise an error if a model does not converge. (Logical)

rm\_nc

Remove non-converged models from output. (Logical)

parallel

Whether to validate the list of models in parallel. (Logical)

verbose

 $Remember \ to \ register \ a \ parallel \ backend \ first. \ E.g. \ with \ doParallel::register DoParallel.$ 

fit and which model function was applied. (Logical)

link, models, model\_verbose

Deprecated.

## **Details**

Packages used:

## **Models:**

Gaussian: stats::lm, lme4::lmer Binomial: stats::glm, lme4::glmer

## **Results:**

Shared:

AIC: stats::AIC AICc: MuMIn::AICc BIC:stats::BIC

Gaussian:

r2m: MuMIn::r.squaredGLMM r2c: MuMIn::r.squaredGLMM

Binomial:

ROC and AUC: pROC::roc

#### Value

tibble with the results and model objects.

#### **Shared across families:**

A nested tibble with **coefficients** of the models from all iterations.

Count of convergence warnings. Consider discarding models that did not converge.

Count of other warnings. These are warnings without keywords such as "convergence".

Count of Singular Fit messages. See lme4::isSingular for more information.

Nested tibble with the warnings and messages caught for each model.

Specified family.

Nested **model** objects.

Name of **dependent** variable.

Names of **fixed** effects.

Names of **random** effects, if any.

Nested tibble with preprocessing parameters, if any.

**Gaussian Results:** 

RMSE, MAE, NRMSE(IQR), RRSE, RAE, RMSLE, AIC, AICc, and BIC.

See the additional metrics (disabled by default) at ?gaussian\_metrics.

A nested tibble with the **predictions** and targets.

#### **Binomial Results:**

Based on predictions of the test set, a confusion matrix and ROC curve are used to get the following: ROC:

AUC, Lower CI, and Upper CI.

Confusion Matrix:

Balanced Accuracy, F1, Sensitivity, Specificity, Positive Predictive Value, Negative Predictive Value, Kappa, Detection Rate, Detection Prevalence, Prevalence, and MCC (Matthews correlation coefficient).

See the additional metrics (disabled by default) at ?binomial\_metrics.

Also includes:

A nested tibble with **predictions**, predicted classes (depends on cutoff), and the targets. Note, that the predictions are *not necessarily* of the *specified* positive class, but of the *model's* positive class (second level of dependent variable, alphabetically).

The pROC::roc ROC curve object(s).

A nested tibble with the **confusion matrix**/matrices. The Pos\_ columns tells you whether a row is a True Positive (TP), True Negative (TN), False Positive (FP), or False Negative (FN), depending on which level is the "positive" class. I.e. the level you wish to predict.

The name of the **Positive Class**.

#### Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

## See Also

Other validation functions: cross\_validate(), cross\_validate\_fn(), validate\_fn()

## **Examples**

```
# Attach packages
library(cvms)
library(groupdata2) # partition()
library(dplyr) # %>% arrange()
# Data is part of cvms
data <- participant.scores</pre>
# Set seed for reproducibility
set.seed(7)
# Partition data
# Keep as single data frame
# We could also have fed validate() separate train and test sets.
data_partitioned <- partition(</pre>
  data,
  p = 0.7,
  cat_col = "diagnosis",
  id_col = "participant",
```

```
list_out = FALSE
) %>%
 arrange(.partitions)
# Validate a model
# Gaussian
validate(
 data_partitioned,
 formulas = "score~diagnosis",
 partitions_col = ".partitions",
 family = "gaussian",
 REML = FALSE
# Binomial
validate(data_partitioned,
 formulas = "diagnosis~score",
 partitions_col = ".partitions",
 family = "binomial"
)
## Feed separate train and test sets
# Partition data to list of data frames
# The first data frame will be train (70% of the data)
# The second will be test (30% of the data)
data_partitioned <- partition(</pre>
 data,
 p = 0.7,
 cat_col = "diagnosis",
 id_col = "participant",
 list_out = TRUE
)
train_data <- data_partitioned[[1]]</pre>
test_data <- data_partitioned[[2]]</pre>
# Validate a model
# Gaussian
validate(
 train_data,
 test_data = test_data,
 formulas = "score~diagnosis",
 family = "gaussian",
 REML = FALSE
)
```

# **Description**

## [Experimental]

Fit your model function on a training set and validate it by predicting a test/validation set. Validate different hyperparameter combinations and formulas at once. Preprocess the train/test split. Returns results and fitted models in a tibble for easy reporting and further analysis.

Compared to validate(), this function allows you supply a custom model function, a predict function, a preprocess function and the hyperparameter values to validate.

Supports regression and classification (binary and multiclass). See `type`.

Note that some metrics may not be computable for some types of model objects.

# Usage

```
validate_fn(
  train_data,
  formulas,
  type,
 model_fn,
 predict_fn,
  test_data = NULL,
  preprocess_fn = NULL,
  preprocess_once = FALSE,
 hyperparameters = NULL,
  partitions_col = ".partitions",
  cutoff = 0.5,
  positive = 2,
 metrics = list(),
  rm_nc = FALSE,
 parallel = FALSE,
  verbose = TRUE
)
```

#### **Arguments**

```
data.frame.

Can contain a grouping factor for identifying partitions - as made with groupdata2::partition().

See `partitions_col`.

Model formulas as strings. (Character)

Will be converted to formula objects before being passed to `model_fn`.

E.g. c("y~x", "y~z").

Can contain random effects.

E.g. c("y~x+(1|r)", "y~z+(1|r)").

type

Type of evaluation to perform:

"gaussian" for regression (like linear regression).

"binomial" for binary classification.

"multinomial" for multiclass classification.
```

model\_fn

Model function that returns a fitted model object. Will usually wrap an existing model function like e1071::svm or nnet::multinom.

Must have the following function arguments:

function(train\_data, formula,

hyperparameters)

predict\_fn

Function for predicting the targets in the test folds/sets using the fitted model object. Will usually wrap stats::predict(), but doesn't have to.

Must have the following function arguments:

Must return predictions in the following formats, depending on `type`:

**Binomial:** vector or one-column matrix / data.frame with probabilities (0-1) of the second class, alphabetically. E.g.:

c(0.3, 0.5, 0.1, 0.5)

N.B. When unsure whether a model type produces probabilities based off the alphabetic order of your classes, using 0 and 1 as classes in the dependent variable instead of the class names should increase the chance of getting probabilities of the right class.

**Gaussian:** vector or one-column matrix / data.frame with the predicted value. E.g.:

c(3.7, 0.9, 1.2, 7.3)

**Multinomial:** data. frame with one column per class containing probabilities of the class. Column names should be identical to how the class names are written in the target column. E.g.:

class_3	class_2	class_1
0.203	0.528	0.269
0.310	0.322	0.368
0.254	0.371	0.375

test\_data

data.frame. If specifying `partitions\_col`, this can be NULL.

preprocess\_fn

Function for preprocessing the training and test sets.

Can, for instance, be used to standardize both the training and test sets with the scaling and centering parameters from the training set.

Must have the following function arguments:

function(train\_data, test\_data,

formula, hyperparameters)

Must return a list with the preprocessed `train\_data` and `test\_data`. It may also contain a tibble with the parameters used in preprocessing:

```
list("train" = train_data,
```

```
"test" = test_data,
```

"parameters" = preprocess\_parameters)

Additional elements in the returned list will be ignored.

The optional parameters tibble will be included in the output. It could have the following format:

Measure	var_1	var_2
Mean	37.921	88.231
SD	12.4	5.986

N.B. When `preprocess\_once` is FALSE, the current formula and hyperparameters will be provided. Otherwise, these arguments will be NULL.

#### preprocess\_once

Whether to apply the preprocessing once (**ignoring** the formula and hyperparameters arguments in `preprocess\_fn`) or for every model separately. (Logical)

When preprocessing does not depend on the current formula or hyperparameters, we can do the preprocessing of each train/test split once, to save time. This may require holding a lot more data in memory though, why it is not the default setting.

#### hyperparameters

Either a named list with hyperparameter values to combine in a grid or a data.frame with one row per hyperparameter combination.

Named list for grid search: Add ".n" to sample the combinations. Can be the number of combinations to use, or a percentage between  $\emptyset$  and 1.

```
list(".n" = 10, # sample 10 combinations
    "lrn_rate" = c(0.1, 0.01, 0.001),
    "h_layers" = c(10, 100, 1000),
    "drop_out" = runif(5, 0.3, 0.7))
```

data.frame with specific hyperparameter combinations: One row per combination to test.

E.g.

lrn_rate	h_layers	drop_out
0.1	10	0.65
0.1	1000	0.65
0.01	1000	0.63

partitions\_col Name of grouping factor for identifying partitions. (Character)

Rows with the value 1 in `partitions\_col` are used as training set and rows with the value 2 are used as test set.

N.B. Only used if 'test\_data' is NULL.

cutoff Threshold for predicted classes. (Numeric)

N.B. Binomial models only

positive Level from dependent variable to predict. Either as character (*preferable*) or level index (1 or 2 - alphabetically).

E.g. if we have the levels "cat" and "dog" and we want "dog" to be the positive class, we can either provide "dog" or 2, as alphabetically, "dog" comes after "cat".

**Note:** For *reproducibility*, it's preferable to **specify the name directly**, as different locales may sort the levels differently.

Used when calculating confusion matrix metrics and creating ROC curves.

The Process column in the output can be used to verify this setting.

N.B. Only affects evaluation metrics, not the model training or returned predictions.

## N.B. Binomial models only.

metrics

list for enabling/disabling metrics.

E.g. list("RMSE" = FALSE) would remove RMSE from the regression results, and list("Accuracy" = TRUE) would add the regular Accuracy metric to the classification results. Default values (TRUE/FALSE) will be used for the remaining available metrics.

You can enable/disable all metrics at once by including "all" = TRUE/FALSE in the list. This is done prior to enabling/disabling individual metrics, why f.i. list("all" = FALSE, "RMSE" = TRUE) would return only the RMSE metric.

The list can be created with gaussian\_metrics(), binomial\_metrics(), or multinomial\_metrics().

Also accepts the string "all".

rm\_nc Remove non-converged models from output. (Logical)

parallel Whether to cross-validate the list of models in parallel. (Logical)

Remember to register a parallel backend first. E.g. with doParallel::registerDoParallel.

verbose Whether to message process information like the number of model instances to

fit. (Logical)

#### **Details**

# Packages used:

#### **Results:**

Shared:

AIC: stats::AIC AICc: MuMIn::AICc BIC: stats::BIC

Gaussian:

r2m: MuMIn::r.squaredGLMM r2c: MuMIn::r.squaredGLMM Binomial and Multinomial: ROC and related metrics:

Binomial: pROC::roc

Multinomial: pROC::multiclass.roc

#### Value

tibble with the results and model objects.

#### **Shared across families:**

A nested tibble with **coefficients** of the models. The coefficients are extracted from the model object with parameters::model\_parameters() or coef() (with some restrictions on the output). If these attempts fail, a default coefficients tibble filled with NAs is returned.

Nested tibble with the used **preprocessing parameters**, if a passed `preprocess\_fn` returns the parameters in a tibble.

Count of **convergence warnings**, using a limited set of keywords (e.g. "convergence"). If a convergence warning does not contain one of these keywords, it will be counted with **other warnings**. Consider discarding models that did not converge on all iterations. Note: you might still see results, but these should be taken with a grain of salt!

Nested tibble with the warnings and messages caught for each model.

Specified family.

Nested **model** objects.

Name of **dependent** variable.

Names of **fixed** effects.

Names of random effects, if any.

**Gaussian Results:** 

RMSE, MAE, NRMSE(IQR), RRSE, RAE, and RMSLE.

See the additional metrics (disabled by default) at ?gaussian\_metrics.

A nested tibble with the **predictions** and targets.

**Binomial Results:** 

Based on predictions of the test set, a confusion matrix and a ROC curve are created to get the following:

ROC:

AUC, Lower CI, and Upper CI

Confusion Matrix:

Balanced Accuracy, F1, Sensitivity, Specificity, Positive Predictive Value, Negative Predictive Value, Kappa, Detection Rate, Detection Prevalence, Prevalence, and MCC (Matthews correlation coefficient).

See the additional metrics (disabled by default) at ?binomial\_metrics.

Also includes:

A nested tibble with **predictions**, predicted classes (depends on cutoff), and the targets. Note, that the predictions are *not necessarily* of the *specified* positive class, but of the *model's* positive class (second level of dependent variable, alphabetically).

The pROC::roc ROC curve object(s).

A nested tibble with the **confusion matrix**/matrices. The Pos\_ columns tells you whether a row is a True Positive (TP), True Negative (TN), False Positive (FP), or False Negative (FN), depending on which level is the "positive" class. I.e. the level you wish to predict.

The name of the **Positive Class**.

#### **Multinomial Results:**

For each class, a *one-vs-all* binomial evaluation is performed. This creates a **Class Level Results** tibble containing the same metrics as the binomial results described above (excluding MCC, AUC, Lower CI and Upper CI), along with a count of the class in the target column (Support). These metrics are used to calculate the **macro-averaged** metrics. The nested class level results tibble is also included in the output tibble, and could be reported along with the macro and overall metrics.

The output tibble contains the macro and overall metrics. The metrics that share their name with the metrics in the nested class level results tibble are averages of those metrics (note: does not remove NAs before averaging). In addition to these, it also includes the Overall Accuracy and the multiclass MCC.

**Note:** Balanced Accuracy is the macro-averaged metric, *not* the macro sensitivity as sometimes used!

Other available metrics (disabled by default, see metrics): Accuracy, *multiclass* AUC, Weighted Balanced Accuracy, Weighted Accuracy, Weighted F1, Weighted Sensitivity, Weighted Sensitivity, Weighted Sensitivity, Weighted Pos Pred Value, Weighted Neg Pred Value, Weighted Kappa, Weighted Detection Rate, Weighted Detection Prevalence, and Weighted Prevalence.

Note that the "Weighted" average metrics are weighted by the Support.

Also includes:

A nested tibble with the **predictions**, predicted classes, and targets.

A list of **ROC** curve objects when AUC is enabled.

A nested tibble with the multiclass **Confusion Matrix**.

#### **Class Level Results**

Besides the binomial evaluation metrics and the Support, the nested class level results tibble also contains a nested tibble with the **Confusion Matrix** from the one-vs-all evaluation. The Pos\_ columns tells you whether a row is a True Positive (TP), True Negative (TN), False Positive (FP), or False Negative (FN), depending on which level is the "positive" class. In our case, 1 is the current class and 0 represents all the other classes together.

#### Author(s)

Ludvig Renbo Olsen, <r-pkgs@ludvigolsen.dk>

## See Also

Other validation functions: cross\_validate(), cross\_validate\_fn(), validate()

# Examples

```
# Attach packages
library(cvms)
library(groupdata2) # fold()
library(dplyr) # %>% arrange() mutate()
```

# Note: More examples of custom functions can be found at:

```
# model_fn: model_functions()
# predict_fn: predict_functions()
# preprocess_fn: preprocess_functions()
# Data is part of cvms
data <- participant.scores</pre>
# Set seed for reproducibility
set.seed(7)
# Fold data
data <- partition(</pre>
 data,
 p = 0.8,
 cat_col = "diagnosis",
 id_col = "participant",
 list_out = FALSE
 mutate(diagnosis = as.factor(diagnosis)) %>%
 arrange(.partitions)
# Formulas to validate
formula_gaussian <- "score ~ diagnosis"</pre>
formula_binomial <- "diagnosis ~ score"</pre>
# Gaussian
# Create model function that returns a fitted model object
lm_model_fn <- function(train_data, formula, hyperparameters) {</pre>
 lm(formula = formula, data = train_data)
}
# Create predict function that returns the predictions
lm_predict_fn <- function(test_data, model, formula,</pre>
                           hyperparameters, train_data) {
 stats::predict(
    object = model,
    newdata = test_data,
    type = "response",
    allow.new.levels = TRUE
 )
}
# Validate the model function
v <- validate_fn(</pre>
 data,
 formulas = formula_gaussian,
 type = "gaussian",
 model_fn = lm_model_fn,
 predict_fn = lm_predict_fn,
```

```
partitions_col = ".partitions"
# Extract model object
v$Model[[1]]
# Binomial
# Create model function that returns a fitted model object
glm_model_fn <- function(train_data, formula, hyperparameters) {</pre>
 glm(formula = formula, data = train_data, family = "binomial")
# Create predict function that returns the predictions
glm_predict_fn <- function(test_data, model, formula,</pre>
                           hyperparameters, train_data) {
 stats::predict(
   object = model,
   newdata = test_data,
   type = "response",
   allow.new.levels = TRUE
 )
}
# Validate the model function
validate_fn(
 data,
 formulas = formula_binomial,
 type = "binomial",
 model_fn = glm_model_fn,
 predict_fn = glm_predict_fn,
 partitions_col = ".partitions"
)
# Support Vector Machine (svm)
# with known hyperparameters
\# Only run if the `e1071` package is installed
if (requireNamespace("e1071", quietly = TRUE)){
# Create model function that returns a fitted model object
# We use the hyperparameters arg to pass in the kernel and cost values
# These will usually have been found with cross_validate_fn()
svm_model_fn <- function(train_data, formula, hyperparameters) {</pre>
 # Expected hyperparameters:
 # - kernel
```

```
# - cost
 if (!"kernel" %in% names(hyperparameters))
    stop("'hyperparameters' must include 'kernel'")
 if (!"cost" %in% names(hyperparameters))
    stop("'hyperparameters' must include 'cost'")
 e1071::svm(
    formula = formula,
    data = train_data,
   kernel = hyperparameters[["kernel"]],
    cost = hyperparameters[["cost"]],
    scale = FALSE,
    type = "C-classification",
    probability = TRUE
 )
}
# Create predict function that returns the predictions
svm_predict_fn <- function(test_data, model, formula,</pre>
                           hyperparameters, train_data) {
 predictions <- stats::predict(</pre>
    object = model,
   newdata = test_data,
   allow.new.levels = TRUE,
   probability = TRUE
 )
 # Extract probabilities
 probabilities <- dplyr::as_tibble(</pre>
    attr(predictions, "probabilities")
 )
 # Return second column
 probabilities[[2]]
}
# Specify hyperparameters to use
# We found these in the examples in ?cross_validate_fn()
svm_hparams <- list(</pre>
 "kernel" = "linear",
  "cost" = 10
)
# Validate the model function
validate_fn(
 data,
 formulas = formula_binomial,
 type = "binomial",
 model_fn = svm_model_fn,
 predict_fn = svm_predict_fn,
 hyperparameters = svm_hparams,
 partitions_col = ".partitions"
)
```

108 wines

} # closes `e1071` package check

wines

Wine varieties

# Description

A list of wine varieties in an approximately Zipfian distribution, ordered by descending frequencies.

## **Format**

A data.frame with 368 rows and 1 variable:

Variety Wine variety, 10 levels

# **Details**

Based on the wine-reviews (v4) kaggle dataset by Zack Thoutt: https://www.kaggle.com/zynicide/wine-reviews

# Author(s)

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