Package 'yardstick'

July 13, 2020

Type Package

Title Tidy Characterizations of Model Performance

Version 0.0.7

Description Tidy tools for quantifying how well model fits to a data set such as confusion matrices, class probability curve summaries, and regression metrics (e.g., RMSE).

License GPL-2

URL <https://github.com/tidymodels/yardstick>,

<https://yardstick.tidymodels.org>

BugReports <https://github.com/tidymodels/yardstick/issues>

Depends R ($>= 2.10$)

Imports dplyr ($>= 0.8.5$), generics, pROC ($>= 1.15.0$), rlang ($>=$ 0.4.0), tidyselect, utils

Suggests covr, crayon, ggplot2, kableExtra, knitr, purrr, rmarkdown, testthat, tidyr

VignetteBuilder knitr

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Repository CRAN

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R topics documented:

accuracy 3

accuracy *Accuracy*

Description

Accuracy is the proportion of the data that are predicted correctly.

Usage

```
accuracy(data, ...)
## S3 method for class 'data.frame'
accuracy(data, truth, estimate, na_rm = TRUE, ...)
```

```
accuracy_vec(truth, estimate, na_rm = TRUE, ...)
```
Arguments

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups. For accuracy_vec(), a single numeric value (or NA).

Multiclass

Accuracy extends naturally to multiclass scenarios. Because of this, macro and micro averaging are not implemented.

Author(s)

Max Kuhn

See Also

```
Other class metrics: bal_accuracy(), detection_prevalence(), f_meas(), j_index(), kap(),
mcc(), npv(), ppv(), precision(), recall(), sens(), spec()
```
Examples

```
library(dplyr)
data("two_class_example")
data("hpc_cv")
# Two class
accuracy(two_class_example, truth, predicted)
# Multiclass
# accuracy() has a natural multiclass extension
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  accuracy(obs, pred)
# Groups are respected
hpc_cv %>%
  group_by(Resample) %>%
  accuracy(obs, pred)
```
average_precision *Area under the precision recall curve*

Description

average_precision() is an alternative to pr_auc() that avoids any ambiguity about what the value of precision should be when $recall == 0$ and there are not yet any false positive values (some say it should be 0, others say 1, others say undefined).

It computes a weighted average of the precision values returned from [pr_curve\(\)](#page-74-1), where the weights are the increase in recall from the previous threshold. See $pr_curve()$ for the full curve.

Usage

```
average_precision(data, ...)
## S3 method for class 'data.frame'
average_precision(
 data,
  truth,
  ...,
```
average_precision 5

```
estimator = NULL,
 na\_rm = TRUE,event_level = yardstick_event_level()
\mathcal{L}average_precision_vec(
 truth,
 estimate,
 estimator = NULL,
 na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
)
```
Arguments

Details

The computation for average precision is a weighted average of the precision values. Assuming you have n rows returned from [pr_curve\(\)](#page-74-1), it is a sum from 2 to n, multiplying the precision value p_i by the increase in recall over the previous threshold, $r_i - r_i(i-1)$.

$$
AP = \sum (r_i - r_{i-1}) \cdot p_i
$$

By summing from 2 to n, the precision value p_1 is never used. While $pr_curve()$ returns a value for p_1 , it is technically undefined as tp / (tp + fp) with tp = 0 and fp = 0. A common convention is to use 1 for p_1, but this metric has the nice property of avoiding the ambiguity. On the other hand, r_1 is well defined as long as there are some events (p), and it is tp / p with tp = 0, so r_1 = Ω .

When p_1 is defined as 1, the average_precision() and roc_auc() values are often very close to one another.

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For average_precision_vec(), a single numeric value (or NA).

Multiclass

Macro and macro-weighted averaging is available for this metric. The default is to select macro averaging if a truth factor with more than 2 levels is provided. Otherwise, a standard binary calculation is done. See vignette("multiclass","yardstick") for more information.

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

See Also

[pr_curve\(\)](#page-74-1) for computing the full precision recall curve.

[pr_auc\(\)](#page-71-1) for computing the area under the precision recall curve using the trapezoidal rule.

Other class probability metrics: [gain_capture\(](#page-19-1)), [mn_log_loss\(](#page-55-1)), [pr_auc\(](#page-71-1)), [roc_auc\(](#page-81-1)), [roc_aunp\(](#page-85-1)), [roc_aunu\(](#page-88-1))

```
# ---------------------------------------------------------------------------
# Two class example
# `truth` is a 2 level factor. The first level is `"Class1"`, which is the
# "event of interest" by default in yardstick. See the Relevant Level
# section above.
data(two_class_example)
# Binary metrics using class probabilities take a factor `truth` column,
```

```
# and a single class probability column containing the probabilities of
# the event of interest. Here, since `"Class1"` is the first level of
```
bal_accuracy 7

```
# `"truth"`, it is the event of interest and we pass in probabilities for it.
average_precision(two_class_example, truth, Class1)
# ---------------------------------------------------------------------------
# Multiclass example
# `obs` is a 4 level factor. The first level is `"VF"`, which is the
# "event of interest" by default in yardstick. See the Relevant Level
# section above.
data(hpc_cv)
# You can use the col1:colN tidyselect syntax
library(dplyr)
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  average_precision(obs, VF:L)
# Change the first level of `obs` from `"VF"` to `"M"` to alter the
# event of interest. The class probability columns should be supplied
# in the same order as the levels.
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  mutate(obs = relevel(obs, "M")) %>%
  average_precision(obs, M, VF:L)
# Groups are respected
hpc_cv %>%
  group_by(Resample) %>%
  average_precision(obs, VF:L)
# Weighted macro averaging
hpc_cv %>%
  group_by(Resample) %>%
  average_precision(obs, VF:L, estimator = "macro_weighted")
# Vector version
# Supply a matrix of class probabilities
fold1 <- hpc_cv %>%
  filter(Resample == "Fold01")
average_precision_vec(
   truth = fold1$obs,
   matrix(
     c(fold1$VF, fold1$F, fold1$M, fold1$L),
     ncol = 4)
)
```


Description

Balanced accuracy is computed here as the average of [sens\(\)](#page-101-1) and [spec\(\)](#page-107-1).

Usage

```
bal_accuracy(data, ...)
## S3 method for class 'data.frame'
bal_accuracy(
 data,
 truth,
 estimate,
 estimator = NULL,
 na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
\mathcal{L}bal_accuracy_vec(
 truth,
 estimate,
 estimator = NULL,
 na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
\mathcal{L}
```
Arguments

"binary". The default uses an internal helper that generally defaults to "first", however, if the deprecated global option yardstick.event_first is set, that will be used instead with a warning.

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For bal_accuracy_vec(), a single numeric value (or NA).

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Multiclass

Macro, micro, and macro-weighted averaging is available for this metric. The default is to select macro averaging if a truth factor with more than 2 levels is provided. Otherwise, a standard binary calculation is done. See vignette("multiclass","yardstick") for more information.

Author(s)

Max Kuhn

See Also

```
Other class metrics: accuracy(), detection_prevalence(), f_meas(), j_index(), kap(), mcc(),
npv(), ppv(), precision(), recall(), sens(), spec()
```
Examples

```
# Two class
data("two_class_example")
bal_accuracy(two_class_example, truth, predicted)
```

```
# Multiclass
library(dplyr)
data(hpc_cv)
```

```
hpc_cv %>%
 filter(Resample == "Fold01") %>%
 bal_accuracy(obs, pred)
```
Groups are respected hpc_cv %>%

```
group_by(Resample) %>%
  bal_accuracy(obs, pred)
# Weighted macro averaging
hpc_cv %>%
  group_by(Resample) %>%
  bal_accuracy(obs, pred, estimator = "macro_weighted")
# Vector version
bal_accuracy_vec(
  two_class_example$truth,
  two_class_example$predicted
)
# Making Class2 the "relevant" level
bal_accuracy_vec(
  two_class_example$truth,
  two_class_example$predicted,
  event_level = "second"
\mathcal{L}
```
ccc *Concordance correlation coefficient*

Description

Calculate the concordance correlation coefficient.

Usage

```
ccc(data, ...)
## S3 method for class 'data.frame'
ccc(data, truth, estimate, bias = FALSE, na_rm = TRUE, ...)
ccc_vec(truth, estimate, bias = FALSE, na_rm = TRUE, ...)
```
Arguments

Details

[ccc\(\)](#page-9-1) is a metric of both consistency/correlation and accuracy, while metrics such as [rmse\(\)](#page-79-1) are strictly for accuracy and metrics such as [rsq\(\)](#page-97-1) are strictly for consistency/correlation

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups. For ccc_vec(), a single numeric value (or NA).

Author(s)

Max Kuhn

References

Lin, L. (1989). A concordance correlation coefficient to evaluate reproducibility. *Biometrics*, 45 $(1), 255 - 268.$

Nickerson, C. (1997). A note on "A concordance correlation coefficient to evaluate reproducibility". *Biometrics*, 53(4), 1503-1507.

See Also

Other numeric metrics: [huber_loss_pseudo\(](#page-30-1)), [huber_loss\(](#page-28-1)), [iic\(](#page-32-1)), [mae\(](#page-41-1)), [mape\(](#page-43-1)), [mase\(](#page-44-1)), [mpe\(](#page-57-1)), [rmse\(](#page-79-1)), [rpd\(](#page-93-1)), [rpiq\(](#page-95-1)), [rsq_trad\(](#page-99-1)), [rsq\(](#page-97-1)), [smape\(](#page-105-1))

Other consistency metrics: [rpd\(](#page-93-1)), [rpiq\(](#page-95-1)), [rsq_trad\(](#page-99-1)), [rsq\(](#page-97-1))

Other accuracy metrics: [huber_loss_pseudo\(](#page-30-1)), [huber_loss\(](#page-28-1)), [iic\(](#page-32-1)), [mae\(](#page-41-1)), [mape\(](#page-43-1)), [mase\(](#page-44-1)), [mpe\(](#page-57-1)), [rmse\(](#page-79-1)), [smape\(](#page-105-1))

```
# Supply truth and predictions as bare column names
ccc(solubility_test, solubility, prediction)
```

```
library(dplyr)
```

```
set.seed(1234)
size <- 100
times <-10# create 10 resamples
```

```
solubility_resampled <- bind_rows(
 replicate(
   n = times,
```
12 conf_mat

```
expr = sample_n(solubility_test, size, replace = TRUE),
    simplify = FALSE
  ),
  .id = "resample"
)
# Compute the metric by group
metric_results <- solubility_resampled %>%
  group_by(resample) %>%
  ccc(solubility, prediction)
metric_results
# Resampled mean estimate
metric_results %>%
  summarise(avg_estimate = mean(.estimate))
```
conf_mat *Confusion Matrix for Categorical Data*

Description

Calculates a cross-tabulation of observed and predicted classes.

Usage

```
conf_mat(data, ...)
## S3 method for class 'data.frame'
conf_mat(data, truth, estimate, dnn = c("Prediction", "Truth"), ...)
## S3 method for class 'conf_mat'
tidy(x, \ldots)autoplot.config_matrix() object, type = "mosaic", ...)
```
Arguments

Details

For [conf_mat\(\)](#page-11-1) objects, a broom tidy() method has been created that collapses the cell counts by cell into a data frame for easy manipulation.

There is also a summary() method that computes various classification metrics at once. See [summary.conf_mat\(\)](#page-111-1)

There is a [ggplot2::autoplot\(\)](#page-0-0) method for quickly visualizing the matrix. Both a heatmap and mosaic type is implemented.

The function requires that the factors have exactly the same levels.

Value

conf_mat() produces an object with class conf_mat. This contains the table and other objects. tidy.conf_mat() generates a tibble with columns name (the cell identifier) and value (the cell count).

When used on a grouped data frame, conf_mat() returns a tibble containing columns for the groups along with conf_mat, a list-column where each element is a conf_mat object.

See Also

[summary.conf_mat\(\)](#page-111-1) for computing a large number of metrics from one confusion matrix.

```
library(dplyr)
data("hpc_cv")
# The confusion matrix from a single assessment set (i.e. fold)
cm < - hpc_cv %>%
 filter(Resample == "Fold01") %>%
 conf_mat(obs, pred)
cm
# Now compute the average confusion matrix across all folds in
# terms of the proportion of the data contained in each cell.
# First get the raw cell counts per fold using the `tidy` method
library(purrr)
library(tidyr)
cells_per_resample <- hpc_cv %>%
 group_by(Resample) %>%
 conf_mat(obs, pred) %>%
 mutate(tidied = map(conf_mat, tidy)) %>%
 unnest(tidied)
```

```
# Get the totals per resample
counts_per_resample <- hpc_cv %>%
  group_by(Resample) %>%
  summarize(total = n()) %>%
  left_join(cells_per_resample, by = "Resample") %>%
  # Compute the proportions
  mutate(prop = value/total) %>%
  group_by(name) %>%
  # Average
  summarize(prop = mean(prop))
counts_per_resample
# Now reshape these into a matrix
mean_cmat <- matrix(counts_per_resample$prop, byrow = TRUE, ncol = 4)
rownames(mean_cmat) <- levels(hpc_cv$obs)
colnames(mean_cmat) <- levels(hpc_cv$obs)
round(mean_cmat, 3)
# The confusion matrix can quickly be visualized using autoplot()
library(ggplot2)
autoplot(cm, type = "mosaic")
autoplot(cm, type = "heatmap")
```
detection_prevalence *Detection prevalence*

Description

Detection prevalence is defined as the number of *predicted* positive events (both true positive and false positive) divided by the total number of predictions.

Usage

```
detection_prevalence(data, ...)
## S3 method for class 'data.frame'
detection_prevalence(
 data,
  truth,
  estimate,
 estimator = NULL,
 na_rm = TRUE,
 event_level = yardstick_event_level(),
  ...
)
```


```
detection_prevalence_vec(
  truth,
 estimate,
  estimator = NULL,
 na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
)
```
Arguments

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups. For detection_prevalence_vec(), a single numeric value (or NA).

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Multiclass

Macro, micro, and macro-weighted averaging is available for this metric. The default is to select macro averaging if a truth factor with more than 2 levels is provided. Otherwise, a standard binary calculation is done. See vignette("multiclass","yardstick") for more information.

Author(s)

Max Kuhn

See Also

```
Other class metrics: accuracy(), bal_accuracy(), f_meas(), j_index(), kap(), mcc(), npv(),
ppv(), precision(), recall(), sens(), spec()
```

```
# Two class
data("two_class_example")
detection_prevalence(two_class_example, truth, predicted)
# Multiclass
library(dplyr)
data(hpc_cv)
hpc_cv %>%
 filter(Resample == "Fold01") %>%
 detection_prevalence(obs, pred)
# Groups are respected
hpc_cv %>%
 group_by(Resample) %>%
 detection_prevalence(obs, pred)
# Weighted macro averaging
hpc_cv %>%
 group_by(Resample) %>%
 detection_prevalence(obs, pred, estimator = "macro_weighted")
# Vector version
detection_prevalence_vec(
 two_class_example$truth,
 two_class_example$predicted
\lambda# Making Class2 the "relevant" level
detection_prevalence_vec(
 two_class_example$truth,
 two_class_example$predicted,
 event_level = "second"
\lambda
```


Description

These functions calculate the f_{meas} of a measurement system for finding relevant documents compared to reference results (the truth regarding relevance). Highly related functions are [recall\(\)](#page-76-1) and [precision\(\)](#page-67-1).

Usage

```
f_meas(data, ...)
## S3 method for class 'data.frame'
f_meas(
  data,
  truth,
  estimate,
 beta = 1,
  estimator = NULL,
 na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
\mathcal{L}f_meas_vec(
  truth,
  estimate,
 beta = 1,
 estimator = NULL,
  na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
)
```
Arguments

Details

The measure "F" is a combination of precision and recall (see below).

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For f_meas_vec(), a single numeric value (or NA).

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Multiclass

Macro, micro, and macro-weighted averaging is available for this metric. The default is to select macro averaging if a truth factor with more than 2 levels is provided. Otherwise, a standard binary calculation is done. See vignette("multiclass","yardstick") for more information.

Implementation

Suppose a 2x2 table with notation:

 f_{max} 19

The formulas used here are:

$$
recall = A/(A + C)
$$

$$
precision = A/(A + B)
$$

$$
F_{meas} = (1 + \beta^2) * precision * recall/((\beta^2 * precision) + recall)
$$

See the references for discussions of the statistics.

Author(s)

Max Kuhn

References

Buckland, M., & Gey, F. (1994). The relationship between Recall and Precision. *Journal of the American Society for Information Science*, 45(1), 12-19.

Powers, D. (2007). Evaluation: From Precision, Recall and F Factor to ROC, Informedness, Markedness and Correlation. Technical Report SIE-07-001, Flinders University

See Also

```
Other class metrics: accuracy(), bal_accuracy(), detection_prevalence(), j_index(), kap(),
mcc(), npv(), ppv(), precision(), recall(), sens(), spec()
```
Other relevance metrics: [precision\(](#page-67-1)), [recall\(](#page-76-1))

```
# Two class
data("two_class_example")
f_meas(two_class_example, truth, predicted)
# Multiclass
library(dplyr)
data(hpc_cv)
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  f_meas(obs, pred)
# Groups are respected
hpc_cv %>%
  group_by(Resample) %>%
  f_meas(obs, pred)
# Weighted macro averaging
hpc_cv %>%
  group_by(Resample) %>%
  f_meas(obs, pred, estimator = "macro_weighted")
```

```
# Vector version
f_meas_vec(
  two_class_example$truth,
  two_class_example$predicted
\mathcal{L}# Making Class2 the "relevant" level
f_meas_vec(
  two_class_example$truth,
  two_class_example$predicted,
  event_level = "second"
\overline{\phantom{a}}
```
gain_capture *Gain capture*

Description

gain_capture() is a measure of performance similar to an AUC calculation, but applied to a gain curve.

Usage

```
gain_capture(data, ...)
## S3 method for class 'data.frame'
gain_capture(
  data,
  truth,
  ...,
  estimator = NULL,
  na\_rm = TRUE,event_level = yardstick_event_level()
\mathcal{L}gain_capture_vec(
  truth,
  estimate,
  estimator = NULL,
  na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
\mathcal{L}
```
Arguments

data A data.frame containing the truth and estimate columns.

Details

gain_capture() calculates the area *under* the gain curve, but *above* the baseline, and then divides that by the area *under* a perfect gain curve, but *above* the baseline. It is meant to represent the amount of potential gain "captured" by the model.

The gain_capture() metric is identical to the *accuracy ratio (AR)*, which is also sometimes called the *gini coefficient*. These two are generally calculated on a cumulative accuracy profile curve, but this is the same as a gain curve. See the Engelmann reference for more information.

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For gain_capture_vec(), a single numeric value (or NA).

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Multiclass

Macro and macro-weighted averaging is available for this metric. The default is to select macro averaging if a truth factor with more than 2 levels is provided. Otherwise, a standard binary calculation is done. See vignette("multiclass","yardstick") for more information.

Author(s)

Max Kuhn

References

Engelmann, Bernd & Hayden, Evelyn & Tasche, Dirk (2003). "Measuring the Discriminative Power of Rating Systems," Discussion Paper Series 2: Banking and Financial Studies 2003,01, Deutsche Bundesbank.

See Also

[gain_curve\(\)](#page-22-1) to compute the full gain curve.

Other class probability metrics: [average_precision\(](#page-3-1)), [mn_log_loss\(](#page-55-1)), [pr_auc\(](#page-71-1)), [roc_auc\(](#page-81-1)), [roc_aunp\(](#page-85-1)), [roc_aunu\(](#page-88-1))

```
# ---------------------------------------------------------------------------
# Two class example
# 'truth' is a 2 level factor. The first level is '"Class1"', which is the
# "event of interest" by default in yardstick. See the Relevant Level
# section above.
data(two_class_example)
# Binary metrics using class probabilities take a factor `truth` column,
# and a single class probability column containing the probabilities of
# the event of interest. Here, since `"Class1"` is the first level of
# `"truth"`, it is the event of interest and we pass in probabilities for it.
gain_capture(two_class_example, truth, Class1)
# ---------------------------------------------------------------------------
# Multiclass example
# `obs` is a 4 level factor. The first level is `"VF"`, which is the
# "event of interest" by default in yardstick. See the Relevant Level
# section above.
data(hpc_cv)
# You can use the col1:colN tidyselect syntax
library(dplyr)
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  gain_capture(obs, VF:L)
```
gain_curve 23

```
# Change the first level of `obs` from `"VF"` to `"M"` to alter the
# event of interest. The class probability columns should be supplied
# in the same order as the levels.
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  mutate(obs = relevel(obs, "M")) %>%
  gain_capture(obs, M, VF:L)
# Groups are respected
hpc_cv %>%
  group_by(Resample) %>%
  gain_capture(obs, VF:L)
# Weighted macro averaging
hpc_cv %>%
  group_by(Resample) %>%
  gain_capture(obs, VF:L, estimator = "macro_weighted")
# Vector version
# Supply a matrix of class probabilities
fold1 <- hpc_cv %>%
  filter(Resample == "Fold01")
gain_capture_vec(
   truth = fold1$obs,
   matrix(
    c(fold1$VF, fold1$F, fold1$M, fold1$L),
     ncol = 4)
\mathcal{L}# ---------------------------------------------------------------------------
# Visualize gain_capture()
# Visually, this represents the area under the black curve, but above the
# 45 degree line, divided by the area of the shaded triangle.
library(ggplot2)
autoplot(gain_curve(two_class_example, truth, Class1))
```
gain_curve *Gain curve*

Description

gain_curve() constructs the full gain curve and returns a tibble. See [gain_capture\(\)](#page-19-1) for the relevant area under the gain curve. Also see [lift_curve\(\)](#page-38-1) for a closely related concept.

Usage

```
gain_curve(data, ...)
## S3 method for class 'data.frame'
gain_curve(
 data,
  truth,
  ...,
 na\_rm = TRUE,event_level = yardstick_event_level()
\mathcal{L}
```

```
autoplot.gain_df(object, ...)
```
Arguments

Details

There is a [ggplot2::autoplot\(\)](#page-0-0) method for quickly visualizing the curve. This works for binary and multiclass output, and also works with grouped data (i.e. from resamples). See the examples.

The greater the area between the gain curve and the baseline, the better the model.

Gain curves are identical to CAP curves (cumulative accuracy profile). See the Engelmann reference for more information on CAP curves.

Value

A tibble with class gain_df or gain_grouped_df having columns:

• .n - The index of the current sample.

- .n_events The index of the current *unique* sample. Values with repeated estimate values are given identical indices in this column.
- .percent_tested The cumulative percentage of values tested.
- .percent_found The cumulative percentage of true results relative to the total number of true results.

Gain and Lift Curves

The motivation behind cumulative gain and lift charts is as a visual method to determine the effectiveness of a model when compared to the results one might expect without a model. As an example, without a model, if you were to advertise to a random 10\ to capture 10\ advertised to your entire customer base. Given a model that predicts which customers are more likely to respond, the hope is that you can more accurately target $10\ \text{>}10\$

The calculation to construct gain curves is as follows:

- 1. truth and estimate are placed in descending order by the estimate values (estimate here is a single column supplied in ...).
- 2. The cumulative number of samples with true results relative to the entire number of true results are found. This is the y-axis in a gain chart.

Multiclass

If a multiclass truth column is provided, a one-vs-all approach will be taken to calculate multiple curves, one per level. In this case, there will be an additional column, .level, identifying the "one" column in the one-vs-all calculation.

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Author(s)

Max Kuhn

References

Engelmann, Bernd & Hayden, Evelyn & Tasche, Dirk (2003). "Measuring the Discriminative Power of Rating Systems," Discussion Paper Series 2: Banking and Financial Studies 2003,01, Deutsche Bundesbank.

See Also

Compute the relevant area under the gain curve with [gain_capture\(\)](#page-19-1). Other curve metrics: [lift_curve\(](#page-38-1)), [pr_curve\(](#page-74-1)), [roc_curve\(](#page-90-1))

Examples

```
# ---------------------------------------------------------------------------
# Two class example
# 'truth' is a 2 level factor. The first level is '"Class1"', which is the
# "event of interest" by default in yardstick. See the Relevant Level
# section above.
data(two_class_example)
# Binary metrics using class probabilities take a factor `truth` column,
# and a single class probability column containing the probabilities of
# the event of interest. Here, since `"Class1"` is the first level of
# `"truth"`, it is the event of interest and we pass in probabilities for it.
gain_curve(two_class_example, truth, Class1)
# ---------------------------------------------------------------------------
# `autoplot()`
library(ggplot2)
library(dplyr)
# Use autoplot to visualize
# The top left hand corner of the grey triangle is a "perfect" gain curve
autoplot(gain_curve(two_class_example, truth, Class1))
# Multiclass one-vs-all approach
# One curve per level
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  gain_curve(obs, VF:L) %>%
  autoplot()
# Same as above, but will all of the resamples
# The resample with the minimum (farthest to the left) "perfect" value is
# used to draw the shaded region
hpc_cv %>%
  group_by(Resample) %>%
  gain_curve(obs, VF:L) %>%
  autoplot()
```
get_weights *Developer helpers*

Description

Helpers to be used alongside [metric_vec_template\(\)](#page-54-1) and [metric_summarizer\(\)](#page-52-1) when creating new metrics. See vignette("custom-metrics","yardstick") for more information.

get_weights 27

Usage

get_weights(data, estimator)

finalize_estimator(x, estimator = NULL, metric_class = "default")

finalize_estimator_internal(metric_dispatcher, x, estimator)

dots_to_estimate(data, ...)

```
validate_estimator(estimator, estimator_override = NULL)
```
Arguments

Weight Calculation

get_weights() accepts a confusion matrix and an estimator of type "macro", "micro", or "macro_weighted" and returns the correct weights. It is useful when creating multiclass metrics.

Estimator Selection

finalize_estimator() is the engine for auto-selection of estimator based on the type of x. Generally x is the truth column. This function is called from the vector method of your metric.

finalize_estimator_internal() is an S3 generic that you should extend for your metric if it does not implement *only* the following estimator types: "binary", "macro", "micro", and "macro_weighted". If your metric does support all of these, the default version of finalize_estimator_internal() will autoselect estimator appropriately. If you need to create a method, it should take the form: finalize_estimator_internal.metric_name. Your method for finalize_estimator_internal() should do two things:

- 1. If estimator is NULL, autoselect the estimator based on the type of x and return a single character for the estimator.
- 2. If estimator is not NULL, validate that it is an allowed estimator for your metric and return it.

If you are using the default for finalize_estimator_internal(), the estimator is selected using the following heuristics:

- 1. If estimator is not NULL, it is validated and returned immediately as no auto-selection is needed.
- 2. If \times is a:
	- factor Then "binary" is returned if it has 2 levels, otherwise "macro" is returned.
	- numeric Then "binary" is returned.
	- table Then "binary" is returned if it has 2 columns, otherwise "macro" is returned. This is useful if you have table methods.
	- matrix Then "macro" is returned.

Dots -> Estimate

dots_to_estimate() is useful with class probability metrics that take ... rather than estimate as an argument. It constructs either a single name if 1 input is provided to ... or it constructs a quosure where the expression constructs a matrix of as many columns as are provided to These are eventually evaluated in the summarise() call in [metric_summarizer\(\)](#page-52-1) and evaluate to either a vector or a matrix for further use in the underlying vector functions.

Estimator Validation

validate_estimator() is called from your metric specific method of finalize_estimator_internal() and ensures that a user provided estimator is of the right format and is one of the allowed values.

See Also

[metric_summarizer\(\)](#page-52-1) [metric_vec_template\(\)](#page-54-1)

hpc_cv *Multiclass Probability Predictions*

Description

Multiclass Probability Predictions

Details

This data frame contains the predicted classes and class probabilities for a linear discriminant analysis model fit to the HPC data set from Kuhn and Johnson (2013). These data are the assessment sets from a 10-fold cross-validation scheme. The data column columns for the true class (obs), the class prediction (pred) and columns for each class probability (columns VF, F, M, and L). Additionally, a column for the resample indicator is included.

huber_loss 29

Value

hpc_cv a data frame

Source

Kuhn, M., Johnson, K. (2013) *Applied Predictive Modeling*, Springer

Examples

data(hpc_cv) str(hpc_cv)

```
# `obs` is a 4 level factor. The first level is `"VF"`, which is the
# "event of interest" by default in yardstick. See the Relevant Level
# section in any classification function (such as `?pr_auc`) to see how
# to change this.
levels(hpc_cv$obs)
```

```
huber_loss Huber loss
```
Description

Calculate the Huber loss, a loss function used in robust regression. This loss function is less sensitive to outliers than [rmse\(\)](#page-79-1). This function is quadratic for small residual values and linear for large residual values.

Usage

```
huber_loss(data, ...)
## S3 method for class 'data.frame'
huber_loss(data, truth, estimate, delta = 1, na_rm = TRUE, ...)
huber_loss_vec(truth, estimate, delta = 1, na_rm = TRUE, \ldots)
```
Arguments

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups. For huber_loss_vec(), a single numeric value (or NA).

Author(s)

James Blair

References

Huber, P. (1964). Robust Estimation of a Location Parameter. *Annals of Statistics*, 53 (1), 73-101.

See Also

```
ccc(huber_loss_pseudo(iic(mae(mape(mase(mpe(),
rmse(), rpd(), rpiq(), rsq_trad(), rsq(), smape()
```
Other accuracy metrics: [ccc\(](#page-9-1)), [huber_loss_pseudo\(](#page-30-1)), [iic\(](#page-32-1)), [mae\(](#page-41-1)), [mape\(](#page-43-1)), [mase\(](#page-44-1)), [mpe\(](#page-57-1)), [rmse\(](#page-79-1)), [smape\(](#page-105-1))

```
# Supply truth and predictions as bare column names
huber_loss(solubility_test, solubility, prediction)
```

```
library(dplyr)
set.seed(1234)
size <- 100
times <- 10
# create 10 resamples
solubility_resampled <- bind_rows(
  replicate(
   n = times,
   expr = sample_n(solubility_test, size, replace = TRUE),
   simplify = FALSE
  ),
  .id = "resample"
)
# Compute the metric by group
metric_results <- solubility_resampled %>%
  group_by(resample) %>%
```
huber_loss_pseudo 31

```
huber_loss(solubility, prediction)
metric_results
# Resampled mean estimate
metric_results %>%
  summarise(avg_estimate = mean(.estimate))
```
huber_loss_pseudo *Psuedo-Huber Loss*

Description

Calculate the Pseudo-Huber Loss, a smooth approximation of [huber_loss\(\)](#page-28-1). Like [huber_loss\(\)](#page-28-1), this is less sensitive to outliers than [rmse\(\)](#page-79-1).

Usage

```
huber_loss_pseudo(data, ...)
```
S3 method for class 'data.frame' huber_loss_pseudo(data, truth, estimate, delta = 1, na_rm = TRUE, ...)

huber_loss_pseudo_vec(truth, estimate, delta = 1, na_rm = TRUE, ...)

Arguments

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For huber_loss_pseudo_vec(), a single numeric value (or NA).

Author(s)

James Blair

References

Huber, P. (1964). Robust Estimation of a Location Parameter. *Annals of Statistics*, 53 (1), 73-101.

Hartley, Richard (2004). Multiple View Geometry in Computer Vision. (Second Edition). Page 619.

See Also

```
Other numeric metrics: ccc(), huber_loss(), iic(), mae(), mape(), mase(), mpe(), rmse(),
rpd(), rpiq(), rsq_trad(), rsq(), smape()
```

```
Other accuracy metrics: ccc(), huber_loss(), iic(), mae(), mape(), mase(), mpe(), rmse(),
smape()
```

```
# Supply truth and predictions as bare column names
huber_loss_pseudo(solubility_test, solubility, prediction)
```

```
library(dplyr)
```

```
set.seed(1234)
size <- 100
times <- 10
# create 10 resamples
solubility_resampled <- bind_rows(
  replicate(
    n = times,
    expr = sample_n(solubility_test, size, replace = TRUE),
    simplify = FALSE
  ),
  .id = "resample"
\mathcal{L}# Compute the metric by group
metric_results <- solubility_resampled %>%
  group_by(resample) %>%
  huber_loss_pseudo(solubility, prediction)
metric_results
# Resampled mean estimate
metric_results %>%
  summarise(avg_estimate = mean(.estimate))
```
Description

Calculate the index of ideality of correlation. This metric has been studied in QSPR/QSAR models as a good criterion for the predictive potential of these models. It is highly dependent on the correlation coefficient as well as the mean absolute error.

Note the application of IIC is useless under two conditions:

- When the negative mean absolute error and positive mean absolute error are both zero.
- When the outliers are symmetric. Since outliers are context dependent, please use your own checks to validate whether this restriction holds and whether the resulting IIC has interpretative value.

The IIC is seen as an alternative to the traditional correlation coefficient and is in the same units as the original data.

Usage

```
iic(data, ...)
## S3 method for class 'data.frame'
iic(data, truth, estimate, na_rm = TRUE, ...)
```

```
iic_vec(truth, estimate, na_rm = TRUE, ...)
```
Arguments

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For iic_vec(), a single numeric value (or NA).

 $\frac{1}{33}$

Author(s)

Joyce Cahoon

References

Toropova, A. and Toropov, A. (2017). "The index of ideality of correlation. A criterion of predictability of QSAR models for skin permeability?" *Science of the Total Environment*. 586: 466- 472.

See Also

```
Other numeric metrics: ccc(), huber_loss_pseudo(), huber_loss(), mae(), mape(), mase(),
mpe(), rmse(), rpd(), rpiq(), rsq_trad(), rsq(), smape()
```

```
Other accuracy metrics: ccc(), huber_loss_pseudo(), huber_loss(), mae(), mape(), mase(),
mpe(), rmse(), smape()
```

```
# Supply truth and predictions as bare column names
iic(solubility_test, solubility, prediction)
library(dplyr)
set.seed(1234)
size <- 100
times <- 10
# create 10 resamples
solubility_resampled <- bind_rows(
 replicate(
   n = times,
   expr = sample_n(solubility_test, size, replace = TRUE),
   simplify = FALSE
  ),
  .id = "resample"
)
# Compute the metric by group
metric_results <- solubility_resampled %>%
  group_by(resample) %>%
  iic(solubility, prediction)
metric_results
```

```
# Resampled mean estimate
metric_results %>%
  summarise(avg_estimate = mean(.estimate))
```
j_index *J-index*

Description

Youden's J statistic is defined as:

```
sens()spec() - 1
```
A related metric is Informedness, see the Details section for the relationship.

Usage

```
j_index(data, ...)
## S3 method for class 'data.frame'
j_index(
 data,
  truth,
 estimate,
 estimator = NULL,
 na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
\mathcal{L}j_index_vec(
  truth,
 estimate,
 estimator = NULL,
 na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
)
```
Arguments

Details

The value of the J-index ranges from $[0, 1]$ and is 1 when there are no false positives and no false negatives.

The binary version of J-index is equivalent to the binary concept of Informedness. Macro-weighted J-index is equivalent to multiclass informedness as defined in Powers, David M W (2011), equation (42).

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For j_index_vec(), a single numeric value (or NA).

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Multiclass

Macro, micro, and macro-weighted averaging is available for this metric. The default is to select macro averaging if a truth factor with more than 2 levels is provided. Otherwise, a standard binary calculation is done. See vignette("multiclass","yardstick") for more information.

Author(s)

Max Kuhn
j_index 37

References

Youden, W.J. (1950). "Index for rating diagnostic tests". Cancer. 3: 32-35.

Powers, David M W (2011). "Evaluation: From Precision, Recall and F-Score to ROC, Informedness, Markedness and Correlation". Journal of Machine Learning Technologies. 2 (1): 37-63.

See Also

```
Other class metrics: accuracy(), bal_accuracy(), detection_prevalence(), f_meas(), kap(),
mcc(), npv(), ppv(), precision(), recall(), sens(), spec()
```

```
# Two class
data("two_class_example")
j_index(two_class_example, truth, predicted)
# Multiclass
library(dplyr)
data(hpc_cv)
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  j_index(obs, pred)
# Groups are respected
hpc_cv %>%
  group_by(Resample) %>%
  j_index(obs, pred)
# Weighted macro averaging
hpc_cv %>%
  group_by(Resample) %>%
  j_index(obs, pred, estimator = "macro_weighted")
# Vector version
j_index_vec(
  two_class_example$truth,
  two_class_example$predicted
)
# Making Class2 the "relevant" level
j_index_vec(
  two_class_example$truth,
  two_class_example$predicted,
  event_level = "second"
\lambda
```


Description

Kappa is a similar measure to [accuracy\(\)](#page-2-0), but is normalized by the accuracy that would be expected by chance alone and is very useful when one or more classes have large frequency distributions.

Usage

```
kap(data, ...)
## S3 method for class 'data.frame'
kap(data, truth, estimate, na_rm = TRUE, ...)
```

```
kap_vec(truth, estimate, na_rm = TRUE, ...)
```
Arguments

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups. For kap_vec(), a single numeric value (or NA).

Multiclass

Kappa extends naturally to multiclass scenarios. Because of this, macro and micro averaging are not implemented.

lift_curve 39

Author(s)

Max Kuhn

References

Cohen, J. (1960). "A coefficient of agreement for nominal scales". *Educational and Psychological Measurement*. 20 (1): 37-46.

See Also

```
Other class metrics: accuracy(), bal_accuracy(), detection_prevalence(), f_meas(), j_index(),
mcc(), npv(), ppv(), precision(), recall(), sens(), spec()
```
Examples

```
library(dplyr)
data("two_class_example")
data("hpc_cv")
```
Two class kap(two_class_example, truth, predicted)

```
# Multiclass
# kap() has a natural multiclass extension
hpc_cv %>%
 filter(Resample == "Fold01") %>%
 kap(obs, pred)
# Groups are respected
hpc_cv %>%
 group_by(Resample) %>%
 kap(obs, pred)
```
lift_curve *Lift curve*

Description

lift_curve() constructs the full lift curve and returns a tibble. See [gain_curve\(\)](#page-22-0) for a closely related concept.

Usage

```
lift_curve(data, ...)
## S3 method for class 'data.frame'
lift_curve(
  data,
```

```
truth,
  ...,
 na_rm = TRUE,
 event_level = yardstick_event_level()
)
```

```
autoplot.lift_df(object, ...)
```
Arguments

Details

There is a [ggplot2::autoplot\(\)](#page-0-0) method for quickly visualizing the curve. This works for binary and multiclass output, and also works with grouped data (i.e. from resamples). See the examples.

Value

A tibble with class lift_df or lift_grouped_df having columns:

- .n The index of the current sample.
- .n_events The index of the current *unique* sample. Values with repeated estimate values are given identical indices in this column.
- .percent_tested The cumulative percentage of values tested.
- .lift First calculate the cumulative percentage of true results relative to the total number of true results. Then divide that by .percent_tested.

lift_curve 41

Gain and Lift Curves

The motivation behind cumulative gain and lift charts is as a visual method to determine the effectiveness of a model when compared to the results one might expect without a model. As an example, without a model, if you were to advertise to a random 10\ to capture 10\ advertised to your entire customer base. Given a model that predicts which customers are more likely to respond, the hope is that you can more accurately target $10\ge 10$

The calculation to construct lift curves is as follows:

- 1. truth and estimate are placed in descending order by the estimate values (estimate here is a single column supplied in ...).
- 2. The cumulative number of samples with true results relative to the entire number of true results are found.
- 3. The cumulative \ to construct the lift value. This ratio represents the factor of improvement over an uninformed model. Values >1 represent a valuable model. This is the y-axis of the lift chart.

Multiclass

If a multiclass truth column is provided, a one-vs-all approach will be taken to calculate multiple curves, one per level. In this case, there will be an additional column, .level, identifying the "one" column in the one-vs-all calculation.

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Author(s)

Max Kuhn

See Also

Other curve metrics: [gain_curve\(](#page-22-0)), [pr_curve\(](#page-74-0)), [roc_curve\(](#page-90-0))

```
# ---------------------------------------------------------------------------
# Two class example
# `truth` is a 2 level factor. The first level is `"Class1"`, which is the
# "event of interest" by default in yardstick. See the Relevant Level
# section above.
data(two_class_example)
```

```
# Binary metrics using class probabilities take a factor `truth` column,
# and a single class probability column containing the probabilities of
# the event of interest. Here, since `"Class1"` is the first level of
# `"truth"`, it is the event of interest and we pass in probabilities for it.
lift_curve(two_class_example, truth, Class1)
# ---------------------------------------------------------------------------
# `autoplot()`
library(ggplot2)
library(dplyr)
# Use autoplot to visualize
autoplot(lift_curve(two_class_example, truth, Class1))
# Multiclass one-vs-all approach
# One curve per level
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  lift_curve(obs, VF:L) %>%
  autoplot()
# Same as above, but will all of the resamples
hpc_cv %>%
  group_by(Resample) %>%
  lift_curve(obs, VF:L) %>%
  autoplot()
```
mae *Mean absolute error*

Description

Calculate the mean absolute error. This metric is in the same units as the original data.

Usage

```
mae(data, ...)
## S3 method for class 'data.frame'
mae(data, truth, estimate, na_rm = TRUE, ...)
```

```
mae_vec(truth, estimate, na_rm = TRUE, ...)
```
Arguments

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For mae_vec(), a single numeric value (or NA).

Author(s)

Max Kuhn

See Also

Other numeric metrics: [ccc\(](#page-9-0)), [huber_loss_pseudo\(](#page-30-0)), [huber_loss\(](#page-28-0)), [iic\(](#page-32-0)), [mape\(](#page-43-0)), [mase\(](#page-44-0)), [mpe\(](#page-57-0)), [rmse\(](#page-79-0)), [rpd\(](#page-93-0)), [rpiq\(](#page-95-0)), [rsq_trad\(](#page-99-0)), [rsq\(](#page-97-0)), [smape\(](#page-105-0))

```
Other accuracy metrics: ccc(), huber_loss_pseudo(), huber_loss(), iic(), mape(), mase(),
mpe(), rmse(), smape()
```

```
# Supply truth and predictions as bare column names
mae(solubility_test, solubility, prediction)
```

```
library(dplyr)
```

```
set.seed(1234)
size <- 100
times <- 10
# create 10 resamples
solubility_resampled <- bind_rows(
  replicate(
   n = times,
   expr = sample_n(solubility_test, size, replace = TRUE),
    simplify = FALSE
  ),
  .id = "resample"
\lambda# Compute the metric by group
metric_results <- solubility_resampled %>%
```
44 mape

```
group_by(resample) %>%
 mae(solubility, prediction)
metric_results
# Resampled mean estimate
metric_results %>%
 summarise(avg_estimate = mean(.estimate))
```
mape *Mean absolute percent error*

Description

Calculate the mean absolute percentage error. This metric is in *relative units*.

Usage

```
mape(data, ...)
## S3 method for class 'data.frame'
mape(data, truth, estimate, na_rm = TRUE, ...)
```
mape_vec(truth, estimate, na_rm = TRUE, ...)

Arguments

Details

Note that a value of Inf is returned for mape() when the observed value is negative.

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values. For grouped data frames, the number of rows returned will be the same as the number of groups. For mape_vec(), a single numeric value (or NA).

mase the contract of the contr

Author(s)

Max Kuhn

See Also

```
Other numeric metrics: ccc(), huber_loss_pseudo(), huber_loss(), iic(), mae(), mase(),
mpe(), rmse(), rpd(), rpiq(), rsq_trad(), rsq(), smape()
ccc(huber_loss_pseudo(huber_loss(iic(mae(mase(),
mpe(), rmse(), smape()
```
Examples

```
# Supply truth and predictions as bare column names
mape(solubility_test, solubility, prediction)
library(dplyr)
set.seed(1234)
size <- 100
times <- 10
# create 10 resamples
solubility_resampled <- bind_rows(
 replicate(
   n = times,
   expr = sample_n(solubility_test, size, replace = TRUE),
   simplify = FALSE
 ),
  .id = "resample"
)
# Compute the metric by group
metric_results <- solubility_resampled %>%
 group_by(resample) %>%
 mape(solubility, prediction)
metric_results
# Resampled mean estimate
metric_results %>%
 summarise(avg_estimate = mean(.estimate))
```
mase *Mean absolute scaled error*

Description

Calculate the mean absolute scaled error. This metric is *scale independent* and *symmetric*. It is generally used for comparing forecast error in time series settings. Due to the time series nature of this metric, it is neccesary to order observations in ascending order by time.

Usage

```
mase(data, ...)
## S3 method for class 'data.frame'
mase(data, truth, estimate, m = 1L, mae_train = NULL, na_rm = TRUE, ...)
```

```
mase_vec(truth, estimate, m = 1L, mae_train = NULL, na_rm = TRUE, ...)
```
Arguments

Details

mase() is different from most numeric metrics. The original implementation of mase() calls for using the *in-sample* naive mean absolute error to compute scaled errors with. It uses this instead of the out-of-sample error because there is a chance that the out-of-sample error cannot be computed when forecasting a very short horizon (i.e. the out of sample size is only 1 or 2). However, yardstick only knows about the out-of-sample truth and estimate values. Because of this, the out-of-sample error is used in the computation by default. If the in-sample naive mean absolute error is required and known, it can be passed through in the mae_train argument and it will be used instead. If the in-sample data is available, the naive mean absolute error can easily be computed with mae(data, truth, lagged_truth).

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For mase_vec(), a single numeric value (or NA).

mase and the contract of the c

Author(s)

Alex Hallam

References

Rob J. Hyndman (2006). ANOTHER LOOK AT FORECAST-ACCURACY METRICS FOR IN-TERMITTENT DEMAND. *Foresight*, 4, 46.

See Also

Other numeric metrics: [ccc\(](#page-9-0)), [huber_loss_pseudo\(](#page-30-0)), [huber_loss\(](#page-28-0)), [iic\(](#page-32-0)), [mae\(](#page-41-0)), [mape\(](#page-43-0)), [mpe\(](#page-57-0)), [rmse\(](#page-79-0)), [rpd\(](#page-93-0)), [rpiq\(](#page-95-0)), [rsq_trad\(](#page-99-0)), [rsq\(](#page-97-0)), [smape\(](#page-105-0))

Other accuracy metrics: [ccc\(](#page-9-0)), [huber_loss_pseudo\(](#page-30-0)), [huber_loss\(](#page-28-0)), [iic\(](#page-32-0)), [mae\(](#page-41-0)), [mape\(](#page-43-0)), [mpe\(](#page-57-0)), [rmse\(](#page-79-0)), [smape\(](#page-105-0))

```
# Supply truth and predictions as bare column names
mase(solubility_test, solubility, prediction)
library(dplyr)
set.seed(1234)
size <- 100
times <-10# create 10 resamples
solubility_resampled <- bind_rows(
  replicate(
   n = times,
   expr = sample_n(solubility_test, size, replace = TRUE),
   simplify = FALSE
  ),
  .id = "resample"
\mathcal{L}# Compute the metric by group
metric_results <- solubility_resampled %>%
  group_by(resample) %>%
  mase(solubility, prediction)
metric_results
# Resampled mean estimate
metric_results %>%
  summarise(avg_estimate = mean(.estimate))
```
Description

Matthews correlation coefficient

Usage

```
mcc(data, ...)
```

```
## S3 method for class 'data.frame'
mcc(data, truth, estimate, na_rm = TRUE, ...)
```
mcc_vec(truth, estimate, na_rm = TRUE, ...)

Arguments

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For mcc_vec(), a single numeric value (or NA).

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

metrics and the contract of th

Multiclass

mcc() has a known multiclass generalization and that is computed automatically if a factor with more than 2 levels is provided. Because of this, no averaging methods are provided.

Author(s)

Max Kuhn

References

Giuseppe, J. (2012). "A Comparison of MCC and CEN Error Measures in Multi-Class Prediction". *PLOS ONE*. Vol 7, Iss 8, e41882.

See Also

```
Other class metrics: accuracy(), ball\_accuracy(), detection\_prevalence(), f\_meas(), j\_index(),kap(), npv(), ppv(), precision(), recall(), sens(), spec()
```
Examples

```
library(dplyr)
data("two_class_example")
data("hpc_cv")
# Two class
mcc(two_class_example, truth, predicted)
# Multiclass
# mcc() has a natural multiclass extension
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  mcc(obs, pred)
# Groups are respected
hpc_cv %>%
```
group_by(Resample) %>% mcc(obs, pred)

metrics *General Function to Estimate Performance*

Description

This function estimates one or more common performance estimates depending on the class of truth (see Value below) and returns them in a three column tibble.

Usage

```
metrics(data, ...)
```

```
## S3 method for class 'data.frame'
metrics(data, truth, estimate, ..., options = list(), na_rm = TRUE)
```
Arguments

Value

A three column tibble.

- When truth is a factor, there are rows for [accuracy\(\)](#page-2-0) and the Kappa statistic ([kap\(\)](#page-37-0)).
- When truth has two levels and 1 column of class probabilities is passed to ..., there are rows for the two class versions of $mn_log_loss()$ and $roc_auc()$.
- When truth has more than two levels and a full set of class probabilities are passed to ..., there are rows for the multiclass version of [mn_log_loss\(\)](#page-55-0) and the Hand Till generalization of [roc_auc\(\)](#page-81-0).
- When truth is numeric, there are rows for [rmse\(\)](#page-79-0), [rsq\(\)](#page-97-0), and [mae\(\)](#page-41-0).

See Also

[metric_set\(\)](#page-50-0)

```
# Accuracy and kappa
metrics(two_class_example, truth, predicted)
```
metric_set 51

```
# Add on multinomal log loss and ROC AUC by specifying class prob columns
metrics(two_class_example, truth, predicted, Class1)
# Regression metrics
metrics(solubility_test, truth = solubility, estimate = prediction)
# Multiclass metrics work, but you cannot specify any averaging
# for roc_auc() besides the default, hand_till. Use the specific function
# if you need more customization
library(dplyr)
hpc_cv %>%
  group_by(Resample) %>%
  metrics(obs, pred, VF:L) %>%
  print(n = 40)
```
metric_set *Combine metric functions*

Description

metric_set() allows you to combine multiple metric functions together into a new function that calculates all of them at once.

Usage

```
metric_set(...)
```
Arguments

... The bare names of the functions to be included in the metric set.

Details

All functions must be either:

- Only numeric metrics
- A mix of class metrics or class prob metrics

For instance, rmse() can be used with mae() because they are numeric metrics, but not with accuracy() because it is a classification metric. But accuracy() can be used with roc_auc().

The returned metric function will have a different argument list depending on whether numeric metrics or a mix of class/prob metrics were passed in.

```
# Numeric metric set signature:
fn(
  data,
  truth,
```

```
estimate,
 na_rm = TRUE,
  ...
\lambda# Class / prob metric set signature:
fn(
  data,
  truth,
  ...,
  estimate,
  estimator = NULL,
 na\_rm = TRUE,event_level = yardstick_event_level()
)
```
When mixing class and class prob metrics, pass in the hard predictions (the factor column) as the named argument estimate, and the soft predictions (the class probability columns) as bare column names or tidyselect selectors to

See Also

[metrics\(\)](#page-48-0)

```
library(dplyr)
# Multiple regression metrics
multi_metric <- metric_set(rmse, rsq, ccc)
# The returned function has arguments:
# fn(data, truth, estimate, na_rm = TRUE, ...)
multi_metric(solubility_test, truth = solubility, estimate = prediction)
# Groups are respected on the new metric function
class_metrics <- metric_set(accuracy, kap)
hpc_cv %>%
  group_by(Resample) %>%
  class_metrics(obs, estimate = pred)
# ---------------------------------------------------------------------------
# If you need to set options for certain metrics,
# do so by wrapping the metric and setting the options inside the wrapper,
# passing along truth and estimate as quoted arguments.
# Then add on the function class of the underlying wrapped function,
# and the direction of optimization.
ccc_with_bias <- function(data, truth, estimate, na_rm = TRUE, ...) {
  ccc(
```

```
data = data,
    truth = !! rlang::enquo(truth),
    estimate = !! rlang::enquo(estimate),
   # set bias = TRUE
   bias = TRUE,na\_rm = na\_rm,
    ...
 )
}
# Use `new_numeric_metric()` to formalize this new metric function
ccc_with_bias <- new_numeric_metric(ccc_with_bias, "maximize")
multi_metric2 <- metric_set(rmse, rsq, ccc_with_bias)
multi_metric2(solubility_test, truth = solubility, estimate = prediction)
# ---------------------------------------------------------------------------
# A class probability example:
# Note that, when given class or class prob functions,
# metric_set() returns a function with signature:
# fn(data, truth, ..., estimate)
# to be able to mix class and class prob metrics.
# You must provide the `estimate` column by explicitly naming
# the argument
class_and_probs_metrics <- metric_set(roc_auc, pr_auc, accuracy)
hpc_cv %>%
  group_by(Resample) %>%
  class_and_probs_metrics(obs, VF:L, estimate = pred)
```
metric_summarizer *Developer function for summarizing new metrics*

Description

metric_summarizer() is useful alongside [metric_vec_template\(\)](#page-54-0) for implementing new custom metrics. metric_summarizer() calls the metric function inside dplyr::summarise(). metric_vec_template() is a generalized function that calls the core implementation of a metric function, and includes a number of checks on the types, lengths, and argument inputs. See vignette("custom-metrics","yardstick") for more information.

Usage

```
metric_summarizer(
 metric_nm,
```

```
metric_fn,
  data,
  truth,
  estimate,
  estimator = NULL,
  na\_rm = TRUE,event_level = NULL,
  ...,
  metric_fn_options = list()
\mathcal{L}
```
Arguments

Details

metric_summarizer() is generally called from the data frame version of your metric function. It knows how to call your metric over grouped data frames and returns a tibble consistent with other metrics.

See Also

[metric_vec_template\(\)](#page-54-0) [finalize_estimator\(\)](#page-25-0) [dots_to_estimate\(\)](#page-25-0)

metric_vec_template *Developer function for calling new metrics*

Description

metric_vec_template() is useful alongside [metric_summarizer\(\)](#page-52-0) for implementing new custom metrics. metric_summarizer() calls the metric function inside dplyr::summarise(). metric_vec_template() is a generalized function that calls the core implementation of a metric function, and includes a number of checks on the types, lengths, and argument inputs.

Usage

```
metric_vec_template(
  metric_impl,
  truth,
  estimate,
  na\_rm = TRUE,cls = "numeric",
  estimator = NULL,
  ...
\mathcal{L}
```
Arguments

... Extra arguments to your core metric function, metric_impl, can technically be passed here, but generally the extra args are added through R's scoping rules because the core metric function is created on the fly when the vector method is called.

Details

metric_vec_template() is called from the vector implementation of your metric. Also defined inside your vector implementation is a separate function performing the core implementation of the metric function. This core function is passed along to metric_vec_template() as metric_impl.

See Also

[metric_summarizer\(\)](#page-52-0) [finalize_estimator\(\)](#page-25-0) [dots_to_estimate\(\)](#page-25-0)

mn_log_loss *Mean log loss*

Description

Compute the logarithmic loss of a classification model.

Usage

```
mn_log_loss(data, ...)
## S3 method for class 'data.frame'
mn_log_loss(
  data,
  truth,
  ...,
 na\_rm = TRUE,sum = FALSE,event_level = yardstick_event_level()
)
mn_log_loss_vec(
  truth,
  estimate,
 na\_rm = TRUE,sum = FALSE,
  event_level = yardstick_event_level(),
  ...
)
```
Arguments

Details

Log loss is a measure of the performance of a classification model. A perfect model has a log loss of 0.

Compared with [accuracy\(\)](#page-2-0), log loss takes into account the uncertainty in the prediction and gives a more detailed view into the actual performance. For example, given two input probabilities of .6 and .9 where both are classified as predicting a positive value, say, "Yes", the accuracy metric would interpret them as having the same value. If the true output is "Yes", log loss penalizes .6 because it is "less sure" of it's result compared to the probability of .9.

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For mn_log_loss_vec(), a single numeric value (or NA).

Multiclass

Log loss has a known multiclass extension, and is simply the sum of the log loss values for each class prediction. Because of this, no averaging types are supported.

Author(s)

Max Kuhn

See Also

Other class probability metrics: [average_precision\(](#page-3-0)), [gain_capture\(](#page-19-0)), [pr_auc\(](#page-71-0)), [roc_auc\(](#page-81-0)), [roc_aunp\(](#page-85-0)), [roc_aunu\(](#page-88-0))

```
# Two class
data("two_class_example")
mn_log_loss(two_class_example, truth, Class1)
# Multiclass
library(dplyr)
data(hpc_cv)
# You can use the col1:colN tidyselect syntax
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  mn_log_loss(obs, VF:L)
# Groups are respected
hpc_cv %>%
  group_by(Resample) %>%
  mn_log_loss(obs, VF:L)
# Vector version
# Supply a matrix of class probabilities
fold1 <- hpc_cv %>%
  filter(Resample == "Fold01")
mn_log_loss_vec(
  truth = fold1$obs,
  matrix(
    c(fold1$VF, fold1$F, fold1$M, fold1$L),
     ncol = 4)
\lambda# Supply `...` with quasiquotation
prob_cols <- levels(two_class_example$truth)
mn_log_loss(two_class_example, truth, Class1)
mn_log_loss(two_class_example, truth, !! prob_cols[1])
```
 mpe 59

Description

Calculate the mean percentage error. This metric is in *relative units*. It can be used as a measure of the estimate's bias.

Note that if *any* truth values are 0, a value of: -Inf (estimate > 0), Inf (estimate < 0), or NaN (estimate $== 0$) is returned for mpe().

Usage

```
mpe(data, ...)
## S3 method for class 'data.frame'
mpe(data, truth, estimate, na_rm = TRUE, ...)
mpe_vec(truth, estimate, na_rm = TRUE, ...)
```
Arguments

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For mpe_vec(), a single numeric value (or NA).

Author(s)

Thomas Bierhance

See Also

Other numeric metrics: [ccc\(](#page-9-0)), [huber_loss_pseudo\(](#page-30-0)), [huber_loss\(](#page-28-0)), [iic\(](#page-32-0)), [mae\(](#page-41-0)), [mape\(](#page-43-0)), [mase\(](#page-44-0)), [rmse\(](#page-79-0)), [rpd\(](#page-93-0)), [rpiq\(](#page-95-0)), [rsq_trad\(](#page-99-0)), [rsq\(](#page-97-0)), [smape\(](#page-105-0))

Other accuracy metrics: [ccc\(](#page-9-0)), [huber_loss_pseudo\(](#page-30-0)), [huber_loss\(](#page-28-0)), [iic\(](#page-32-0)), [mae\(](#page-41-0)), [mape\(](#page-43-0)), [mase\(](#page-44-0)), [rmse\(](#page-79-0)), [smape\(](#page-105-0))

Examples

```
# `solubility_test$solubility` has zero values with corresponding
# `$prediction` values that are negative. By definition, this causes `Inf`
# to be returned from `mpe()`.
solubility_test[solubility_test$solubility == 0,]
mpe(solubility_test, solubility, prediction)
# We'll remove the zero values for demonstration
solubility_test <- solubility_test[solubility_test$solubility != 0,]
# Supply truth and predictions as bare column names
mpe(solubility_test, solubility, prediction)
library(dplyr)
set.seed(1234)
size <- 100
times <-10# create 10 resamples
solubility_resampled <- bind_rows(
  replicate(
   n = times,
   expr = sample_n(solubility_test, size, replace = TRUE),
   simplify = FALSE
  ),
  .id = "resample"
\mathcal{L}# Compute the metric by group
metric_results <- solubility_resampled %>%
  group_by(resample) %>%
  mpe(solubility, prediction)
metric_results
# Resampled mean estimate
metric_results %>%
  summarise(avg_estimate = mean(.estimate))
```
new-metric *Construct a new metric function*

Description

These functions provide convenient wrappers to create the three types of metric functions in yardstick: numeric metrics, class metrics, and class probability metrics. They add a metric-specific class to fn and attach a direction attribute. These features are used by [metric_set\(\)](#page-50-0) and by [tune](https://tune.tidymodels.org/) when model tuning.

See vignette("custom-metrics") for more information about creating custom metrics.

Usage

```
new_class_metric(fn, direction)
new_prob_metric(fn, direction)
```

```
new_numeric_metric(fn, direction)
```
Arguments

npv *Negative predictive value*

Description

These functions calculate the [npv\(\)](#page-60-0) (negative predictive value) of a measurement system compared to a reference result (the "truth" or gold standard). Highly related functions are [spec\(\)](#page-107-0), [sens\(\)](#page-101-0), and [ppv\(\)](#page-64-0).

Usage

```
npv(data, ...)
## S3 method for class 'data.frame'
npv(
 data,
  truth,
  estimate,
 prevalence = NULL,
  estimator = NULL,
 na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
)
npv_vec(
  truth,
```

```
estimate,
 prevalence = NULL,
 estimator = NULL,
 na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
\mathcal{L}
```
Arguments

Details

The positive predictive value $(ppv()$ $(ppv()$ is defined as the percent of predicted positives that are actually positive while the negative predictive value ([npv\(\)](#page-60-0)) is defined as the percent of negative positives that are actually negative.

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For npv_vec(), a single numeric value (or NA).

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Multiclass

Macro, micro, and macro-weighted averaging is available for this metric. The default is to select macro averaging if a truth factor with more than 2 levels is provided. Otherwise, a standard binary calculation is done. See vignette("multiclass","yardstick") for more information.

Implementation

Suppose a 2x2 table with notation:

The formulas used here are:

Sensitivity =
$$
A/(A + C)
$$

\nSpecificity = $D/(B + D)$
\nPrevalence = $(A + C)/(A + B + C + D)$

P P V = (Sensitivity∗P revalence)/((Sensitivity∗P revalence)+((1−Specif icity)∗(1−P revalence))) $NPV = (Specificity * (1-Prevalence)) / (((1-Sensitivity) * Prevalence) + ((Specificity) * (1-Prevalence)))$

See the references for discussions of the statistics.

Author(s)

Max Kuhn

References

Altman, D.G., Bland, J.M. (1994) "Diagnostic tests 2: predictive values," *British Medical Journal*, vol 309, 102.

See Also

```
Other class metrics: accuracy(), bal_accuracy(), detection_prevalence(), f_meas(), j_index(),
kap(), mcc(), ppv(), precision(), recall(), sens(), spec()
Other sensitivity metrics: ppv(), sens(), spec()
```
 $n_{\rm PV}$ 63

64 pathology

Examples

```
# Two class
data("two_class_example")
npv(two_class_example, truth, predicted)
# Multiclass
library(dplyr)
data(hpc_cv)
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  npv(obs, pred)
# Groups are respected
hpc_cv %>%
  group_by(Resample) %>%
  npv(obs, pred)
# Weighted macro averaging
hpc_cv %>%
  group_by(Resample) %>%
  npv(obs, pred, estimator = "macro_weighted")
# Vector version
npv_vec(
  two_class_example$truth,
  two_class_example$predicted
)
# Making Class2 the "relevant" level
npv_vec(
  two_class_example$truth,
  two_class_example$predicted,
  event_level = "second"
\mathcal{L}
```
pathology *Liver Pathology Data*

Description

Liver Pathology Data

Details

These data have the results of a *x*-ray examination to determine whether liver is abnormal or not (in the scan column) versus the more extensive pathology results that approximate the truth (in pathology).

 ppv 65

Value

pathology a data frame

Source

Altman, D.G., Bland, J.M. (1994) "Diagnostic tests 1: sensitivity and specificity," *British Medical Journal*, vol 308, 1552.

Examples

data(pathology) str(pathology)

ppv *Positive predictive value*

Description

These functions calculate the [ppv\(\)](#page-64-0) (positive predictive value) of a measurement system compared to a reference result (the "truth" or gold standard). Highly related functions are [spec\(\)](#page-107-0), [sens\(\)](#page-101-0), and [npv\(\)](#page-60-0).

Usage

```
ppv(data, ...)
## S3 method for class 'data.frame'
ppv(
  data,
  truth,
  estimate,
 prevalence = NULL,
 estimator = NULL,
 na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
)
ppv_vec(
  truth,
  estimate,
 prevalence = NULL,
 estimator = NULL,
 na_rm = TRUE,
 event_level = yardstick_event_level(),
  ...
\mathcal{L}
```
Arguments

Details

The positive predictive value $(ppv()$ $(ppv()$ is defined as the percent of predicted positives that are actually positive while the negative predictive value $(npv())$ $(npv())$ $(npv())$ is defined as the percent of negative positives that are actually negative.

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For ppv_vec(), a single numeric value (or NA).

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Multiclass

Macro, micro, and macro-weighted averaging is available for this metric. The default is to select macro averaging if a truth factor with more than 2 levels is provided. Otherwise, a standard binary calculation is done. See vignette("multiclass","yardstick") for more information.

Implementation

Suppose a 2x2 table with notation:

The formulas used here are:

 $Sensitivity = A/(A+C)$ $Specificity = D/(B+D)$

$$
Prevalence = (A+C)/(A+B+C+D)
$$

P P V = (Sensitivity∗P revalence)/((Sensitivity∗P revalence)+((1−Specif icity)∗(1−P revalence)))

 $NPV = (Specificity * (1-Prevalence)) / (((1-Sensitivity) * Prevalence) + ((Specificity) * (1-Prevalence)))$

See the references for discussions of the statistics.

Author(s)

Max Kuhn

References

Altman, D.G., Bland, J.M. (1994) "Diagnostic tests 2: predictive values," *British Medical Journal*, vol 309, 102.

See Also

```
Other class metrics: accuracy(), bal_accuracy(), detection_prevalence(), f_meas(), j_index(),
kap(), mcc(), npv(), precision(), recall(), sens(), spec()
```
Other sensitivity metrics: [npv\(](#page-60-0)), [sens\(](#page-101-0)), [spec\(](#page-107-0))

ppv 67

68 precision

Examples

```
# Two class
data("two_class_example")
ppv(two_class_example, truth, predicted)
# Multiclass
library(dplyr)
data(hpc_cv)
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  ppv(obs, pred)
# Groups are respected
hpc_cv %>%
  group_by(Resample) %>%
  ppv(obs, pred)
# Weighted macro averaging
hpc_cv %>%
  group_by(Resample) %>%
  ppv(obs, pred, estimator = "macro_weighted")
# Vector version
ppv_vec(
  two_class_example$truth,
  two_class_example$predicted
\lambda# Making Class2 the "relevant" level
ppv_vec(
  two_class_example$truth,
  two_class_example$predicted,
  event_level = "second"
)
# But what if we think that Class 1 only occurs 40% of the time?
ppv(two_class_example, truth, predicted, prevalence = 0.40)
```
precision *Precision*

Description

These functions calculate the [precision\(\)](#page-67-0) of a measurement system for finding relevant documents compared to reference results (the truth regarding relevance). Highly related functions are [recall\(\)](#page-76-0) and [f_meas\(\)](#page-16-0).

precision 69

Usage

```
precision(data, ...)
## S3 method for class 'data.frame'
precision(
  data,
  truth,
  estimate,
  estimator = NULL,
  na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
\mathcal{L}precision_vec(
  truth,
  estimate,
  estimator = NULL,
  na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
\mathcal{L}
```
Arguments

Details

The precision is the percentage of predicted truly relevant results of the total number of predicted relevant results and characterizes the "purity in retrieval performance" (Buckland and Gey, 1994).

When the denominator of the calculation is θ , precision is undefined. This happens when both # true_positive = 0 and # false_positive = 0 are true, which mean that there were no predicted events. When computing binary precision, a NA value will be returned with a warning. When computing multiclass precision, the individual NA values will be removed, and the computation will procede, with a warning.

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For precision_vec(), a single numeric value (or NA).

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Multiclass

Macro, micro, and macro-weighted averaging is available for this metric. The default is to select macro averaging if a truth factor with more than 2 levels is provided. Otherwise, a standard binary calculation is done. See vignette("multiclass","yardstick") for more information.

Implementation

Suppose a 2x2 table with notation:

The formulas used here are:

$$
recall = A/(A + C)
$$

$$
precision = A/(A + B)
$$

$$
F_{meas} = (1 + \beta^2) * precision * recall/((\beta^2 * precision) + recall)
$$

See the references for discussions of the statistics.

precision 71

Author(s)

Max Kuhn

References

Buckland, M., & Gey, F. (1994). The relationship between Recall and Precision. *Journal of the American Society for Information Science*, 45(1), 12-19.

Powers, D. (2007). Evaluation: From Precision, Recall and F Factor to ROC, Informedness, Markedness and Correlation. Technical Report SIE-07-001, Flinders University

See Also

```
Other class metrics: accuracy(), bal_accuracy(), detection_prevalence(), f_meas(), j_index(),
kap(), mcc(), npv(), ppv(), recall(), sens(), spec()
```

```
Other relevance metrics: f_meas(), recall()
```

```
# Two class
data("two_class_example")
precision(two_class_example, truth, predicted)
# Multiclass
library(dplyr)
data(hpc_cv)
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  precision(obs, pred)
# Groups are respected
hpc_cv %>%
  group_by(Resample) %>%
  precision(obs, pred)
# Weighted macro averaging
hpc_cv %>%
  group_by(Resample) %>%
  precision(obs, pred, estimator = "macro_weighted")
# Vector version
precision_vec(
  two_class_example$truth,
  two_class_example$predicted
\mathcal{L}# Making Class2 the "relevant" level
precision_vec(
  two_class_example$truth,
  two_class_example$predicted,
  event_level = "second"
```
 λ

pr_auc *Area under the precision recall curve*

Description

pr_auc() is a metric that computes the area under the precision recall curve. See [pr_curve\(\)](#page-74-0) for the full curve.

Usage

```
pr_auc(data, ...)
## S3 method for class 'data.frame'
pr_auc(
 data,
  truth,
  ...,
 estimator = NULL,
 na\_rm = TRUE,event_level = yardstick_event_level()
\mathcal{L}pr_auc_vec(
  truth,
 estimate,
 estimator = NULL,
 na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
\mathcal{L}
```
Arguments

pr_2 auc $\hspace{1cm}$ 73

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For pr_auc_vec(), a single numeric value (or NA).

Multiclass

Macro and macro-weighted averaging is available for this metric. The default is to select macro averaging if a truth factor with more than 2 levels is provided. Otherwise, a standard binary calculation is done. See vignette("multiclass","yardstick") for more information.

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Author(s)

Max Kuhn

See Also

[pr_curve\(\)](#page-74-0) for computing the full precision recall curve.

Other class probability metrics: [average_precision\(](#page-3-0)), [gain_capture\(](#page-19-0)), [mn_log_loss\(](#page-55-0)), [roc_auc\(](#page-81-0)), [roc_aunp\(](#page-85-0)), [roc_aunu\(](#page-88-0))

Examples

```
# ---------------------------------------------------------------------------
# Two class example
# 'truth' is a 2 level factor. The first level is '"Class1"', which is the
# "event of interest" by default in yardstick. See the Relevant Level
# section above.
data(two_class_example)
# Binary metrics using class probabilities take a factor `truth` column,
# and a single class probability column containing the probabilities of
# the event of interest. Here, since `"Class1"` is the first level of
# `"truth"`, it is the event of interest and we pass in probabilities for it.
pr_auc(two_class_example, truth, Class1)
# ---------------------------------------------------------------------------
# Multiclass example
# `obs` is a 4 level factor. The first level is `"VF"`, which is the
# "event of interest" by default in yardstick. See the Relevant Level
# section above.
data(hpc_cv)
# You can use the col1:colN tidyselect syntax
library(dplyr)
hpc_cv %>%
 filter(Resample == "Fold01") %>%
  pr_auc(obs, VF:L)
# Change the first level of `obs` from `"VF"` to `"M"` to alter the
# event of interest. The class probability columns should be supplied
# in the same order as the levels.
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  mutate(obs = relevel(obs, "M")) %>%
  pr_auc(obs, M, VF:L)
# Groups are respected
hpc_cv %>%
  group_by(Resample) %>%
  pr_auc(obs, VF:L)
# Weighted macro averaging
hpc_cv %>%
  group_by(Resample) %>%
  pr_auc(obs, VF:L, estimator = "macro_weighted")
# Vector version
# Supply a matrix of class probabilities
fold1 <- hpc_cv %>%
  filter(Resample == "Fold01")
```
pr_curve 75

```
pr_auc_vec(
   truth = fold1$obs,
   matrix(
    c(fold1$VF, fold1$F, fold1$M, fold1$L),
     ncol = 4)
\mathcal{L}
```
pr_curve *Precision recall curve*

Description

pr_curve() constructs the full precision recall curve and returns a tibble. See [pr_auc\(\)](#page-71-0) for the area under the precision recall curve.

Usage

```
pr_curve(data, ...)
## S3 method for class 'data.frame'
pr_curve(data, truth, ..., na_rm = TRUE, event_level = yardstick_event_level())
```
autoplot.pr_df(object, ...)

Arguments

Details

pr_curve() computes the precision at every unique value of the probability column (in addition to infinity).

There is a [ggplot2::autoplot\(\)](#page-0-0) method for quickly visualizing the curve. This works for binary and multiclass output, and also works with grouped data (i.e. from resamples). See the examples.

Value

A tibble with class pr_df or pr_grouped_df having columns .threshold, recall, and precision.

Multiclass

If a multiclass truth column is provided, a one-vs-all approach will be taken to calculate multiple curves, one per level. In this case, there will be an additional column, .level, identifying the "one" column in the one-vs-all calculation.

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Author(s)

Max Kuhn

See Also

Compute the area under the precision recall curve with [pr_auc\(\)](#page-71-0).

Other curve metrics: [gain_curve\(](#page-22-0)), [lift_curve\(](#page-38-0)), [roc_curve\(](#page-90-0))

Examples

```
# ---------------------------------------------------------------------------
# Two class example
# `truth` is a 2 level factor. The first level is `"Class1"`, which is the
# "event of interest" by default in yardstick. See the Relevant Level
# section above.
data(two_class_example)
# Binary metrics using class probabilities take a factor `truth` column,
# and a single class probability column containing the probabilities of
# the event of interest. Here, since `"Class1"` is the first level of
# `"truth"`, it is the event of interest and we pass in probabilities for it.
```
recall and the contract of the

```
# ---------------------------------------------------------------------------
# `autoplot()`
# Visualize the curve using ggplot2 manually
library(ggplot2)
library(dplyr)
pr_curve(two_class_example, truth, Class1) %>%
  ggplot(aes(x = recall, y = precision)) +geom_path() +
  coord_equal() +
  theme_bw()
# Or use autoplot
autoplot(pr_curve(two_class_example, truth, Class1))
# Multiclass one-vs-all approach
# One curve per level
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  pr_curve(obs, VF:L) %>%
  autoplot()
# Same as above, but will all of the resamples
hpc_cv %>%
  group_by(Resample) %>%
  pr_curve(obs, VF:L) %>%
  autoplot()
```
recall *Recall*

Description

These functions calculate the [recall\(\)](#page-76-0) of a measurement system for finding relevant documents compared to reference results (the truth regarding relevance). Highly related functions are [precision\(\)](#page-67-0) and [f_meas\(\)](#page-16-0).

Usage

```
recall(data, ...)
## S3 method for class 'data.frame'
recall(
 data,
  truth,
 estimate,
  estimator = NULL,
 na\_rm = TRUE,
```
78 recall

```
event_level = yardstick_event_level(),
  ...
\mathcal{L}recall_vec(
  truth,
  estimate,
  estimator = NULL,
  na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
)
```
Arguments

Details

The recall (aka sensitivity) is defined as the proportion of relevant results out of the number of samples which were actually relevant. When there are no relevant results, recall is not defined and a value of NA is returned.

When the denominator of the calculation is θ , recall is undefined. This happens when both # true_positive = 0 and $#$ false_negative = 0 are true, which mean that there were no true events. When computing binary recall, a NA value will be returned with a warning. When computing multiclass recall, the individual NA values will be removed, and the computation will procede, with a warning.

recall and the contract of the

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For recall_vec(), a single numeric value (or NA).

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Multiclass

Macro, micro, and macro-weighted averaging is available for this metric. The default is to select macro averaging if a truth factor with more than 2 levels is provided. Otherwise, a standard binary calculation is done. See vignette("multiclass","yardstick") for more information.

Implementation

Suppose a 2x2 table with notation:

The formulas used here are:

$$
recall = A/(A + C)
$$

$$
precision = A/(A + B)
$$

$$
F_{meas} = (1 + \beta^2) * precision * recall/((\beta^2 * precision) + recall)
$$

See the references for discussions of the statistics.

Author(s)

Max Kuhn

References

Buckland, M., & Gey, F. (1994). The relationship between Recall and Precision. *Journal of the American Society for Information Science*, 45(1), 12-19.

Powers, D. (2007). Evaluation: From Precision, Recall and F Factor to ROC, Informedness, Markedness and Correlation. Technical Report SIE-07-001, Flinders University

See Also

```
Other class metrics: accuracy(), ball\_accuracy(), detection\_prevalence(), f\_meas(), j\_index(),kap(), mcc(), npv(), ppv(), precision(), sens(), spec()
```

```
Other relevance metrics: f_meas(), precision()
```
Examples

```
# Two class
data("two_class_example")
recall(two_class_example, truth, predicted)
# Multiclass
library(dplyr)
data(hpc_cv)
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  recall(obs, pred)
# Groups are respected
hpc_cv %>%
  group_by(Resample) %>%
  recall(obs, pred)
# Weighted macro averaging
hpc_cv %>%
  group_by(Resample) %>%
  recall(obs, pred, estimator = "macro_weighted")
# Vector version
recall_vec(
  two_class_example$truth,
  two_class_example$predicted
)
# Making Class2 the "relevant" level
recall_vec(
  two_class_example$truth,
  two_class_example$predicted,
  event_level = "second"
)
```
rmse *Root mean squared error*

Description

Calculate the root mean squared error. rmse() is a metric that is in the same units as the original data.

rmse aangebruik is de staat 1976 en 19

Usage

```
rmse(data, ...)
## S3 method for class 'data.frame'
rmse(data, truth, estimate, na_rm = TRUE, ...)
```

```
rmse_vec(truth, estimate, na_rm = TRUE, ...)
```
Arguments

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For rmse_vec(), a single numeric value (or NA).

Author(s)

Max Kuhn

See Also

Other numeric metrics: [ccc\(](#page-9-0)), [huber_loss_pseudo\(](#page-30-0)), [huber_loss\(](#page-28-0)), [iic\(](#page-32-0)), [mae\(](#page-41-0)), [mape\(](#page-43-0)), [mase\(](#page-44-0)), [mpe\(](#page-57-0)), [rpd\(](#page-93-0)), [rpiq\(](#page-95-0)), [rsq_trad\(](#page-99-0)), [rsq\(](#page-97-0)), [smape\(](#page-105-0))

Other accuracy metrics: [ccc\(](#page-9-0)), [huber_loss_pseudo\(](#page-30-0)), [huber_loss\(](#page-28-0)), [iic\(](#page-32-0)), [mae\(](#page-41-0)), [mape\(](#page-43-0)), [mase\(](#page-44-0)), [mpe\(](#page-57-0)), [smape\(](#page-105-0))

Examples

Supply truth and predictions as bare column names rmse(solubility_test, solubility, prediction)

library(dplyr)

set.seed(1234)

```
size <- 100
times <-10# create 10 resamples
solubility_resampled <- bind_rows(
  replicate(
   n = times,
   expr = sample_n(solubility_test, size, replace = TRUE),
   simplify = FALSE
  ),
  .id = "resample"
\mathcal{L}# Compute the metric by group
metric_results <- solubility_resampled %>%
  group_by(resample) %>%
  rmse(solubility, prediction)
metric_results
# Resampled mean estimate
metric_results %>%
  summarise(avg_estimate = mean(.estimate))
```


```
roc_auc Area under the receiver operator curve
```
Description

roc_auc() is a metric that computes the area under the ROC curve. See [roc_curve\(\)](#page-90-0) for the full curve.

Usage

```
roc_auc(data, ...)
## S3 method for class 'data.frame'
roc_auc(
 data,
  truth,
  ...,
 options = list(),estimator = NULL,
 na_rm = TRUE,
 event_level = yardstick_event_level()
)
roc_auc_vec(
  truth,
```
roc_auc 83

```
estimate,
 options = list(),
 estimator = NULL,
 na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
)
```
Arguments

Details

The underlying direction option in $pROC::roc()$ is forced to direction = "<". This computes the ROC curve assuming that the estimate values are the probability that the "event" occurred, which is what they are always assumed to be in yardstick.

Generally, an ROC AUC value is between 0.5 and 1, with 1 being a perfect prediction model. If your value is between 0 and 0.5, then this implies that you have meaningful information in your model, but it is being applied incorrectly because doing the opposite of what the model predicts would result in an AUC >0.5.

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For roc_auc_vec(), a single numeric value (or NA).

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Multiclass

The default multiclass method for computing roc_auc() is to use the method from Hand, Till, (2001). Unlike macro-averaging, this method is insensitive to class distributions like the binary ROC AUC case. Additionally, while other multiclass techniques will return NA if any levels in truth occur zero times in the actual data, the Hand-Till method will simply ignore those levels in the averaging calculation, with a warning.

Macro and macro-weighted averaging are still provided, even though they are not the default. In fact, macro-weighted averaging corresponds to the same definition of multiclass AUC given by Provost and Domingos (2001).

Author(s)

Max Kuhn

References

Hand, Till (2001). "A Simple Generalisation of the Area Under the ROC Curve for Multiple Class Classification Problems". *Machine Learning*. Vol 45, Iss 2, pp 171-186.

Fawcett (2005). "An introduction to ROC analysis". *Pattern Recognition Letters*. 27 (2006), pp 861-874.

Provost, F., Domingos, P., 2001. "Well-trained PETs: Improving probability estimation trees", CeDER Working Paper #IS-00-04, Stern School of Business, New York University, NY, NY 10012.

See Also

[roc_curve\(\)](#page-90-0) for computing the full ROC curve.

Other class probability metrics: [average_precision\(](#page-3-0)), [gain_capture\(](#page-19-0)), [mn_log_loss\(](#page-55-0)), [pr_auc\(](#page-71-0)), [roc_aunp\(](#page-85-0)), [roc_aunu\(](#page-88-0))

roc_auc 85

Examples

```
# ---------------------------------------------------------------------------
# Two class example
# 'truth' is a 2 level factor. The first level is '"Class1"', which is the
# "event of interest" by default in yardstick. See the Relevant Level
# section above.
data(two_class_example)
# Binary metrics using class probabilities take a factor `truth` column,
# and a single class probability column containing the probabilities of
# the event of interest. Here, since `"Class1"` is the first level of
# `"truth"`, it is the event of interest and we pass in probabilities for it.
roc_auc(two_class_example, truth, Class1)
# ---------------------------------------------------------------------------
# Multiclass example
# `obs` is a 4 level factor. The first level is `"VF"`, which is the
# "event of interest" by default in yardstick. See the Relevant Level
# section above.
data(hpc_cv)
# You can use the col1:colN tidyselect syntax
library(dplyr)
hpc_cv %>%
 filter(Resample == "Fold01") %>%
  roc_auc(obs, VF:L)
# Change the first level of `obs` from `"VF"` to `"M"` to alter the
# event of interest. The class probability columns should be supplied
# in the same order as the levels.
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  mutate(obs = relevel(obs, "M")) %>%
  roc_auc(obs, M, VF:L)
# Groups are respected
hpc_cv %>%
  group_by(Resample) %>%
  roc_auc(obs, VF:L)
# Weighted macro averaging
hpc_cv %>%
  group_by(Resample) %>%
  roc_auc(obs, VF:L, estimator = "macro_weighted")
# Vector version
# Supply a matrix of class probabilities
fold1 <- hpc_cv %>%
  filter(Resample == "Fold01")
```

```
roc_auc_vec(
  truth = fold1$obs,
  matrix(
    c(fold1$VF, fold1$F, fold1$M, fold1$L),
    ncol = 4)
)
# ---------------------------------------------------------------------------
# Options for `pROC::roc()`
# Pass options via a named list and not through `...`!
roc_auc(
 two_class_example,
 truth = truth,
 Class1,
 options = list(smooth = TRUE)
)
```
roc_aunp *Area under the ROC curve of each class against the rest, using the a priori class distribution*

Description

roc_aunp() is a multiclass metric that computes the area under the ROC curve of each class against the rest, using the a priori class distribution. This is equivalent to roc_auc(estimator = "macro_weighted").

Usage

```
roc_aunp(data, ...)
## S3 method for class 'data.frame'
roc_aunp(data, truth, ..., options = list(), na_rm = TRUE)
roc_aunp_vec(truth, estimate, options = list(), na_rm = TRUE, ...)
```
Arguments

roc_aunp 87

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For roc_aunp_vec(), a single numeric value (or NA).

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Multiclass

This multiclass method for computing the area under the ROC curve uses the a priori class distribution and is equivalent to roc_auc(estimator = "macro_weighted").

Author(s)

Julia Silge

References

Ferri, C., Hernández-Orallo, J., & Modroiu, R. (2009). "An experimental comparison of performance measures for classification". *Pattern Recognition Letters*. 30 (1), pp 27-38.

See Also

[roc_aunu\(\)](#page-88-0) for computing the area under the ROC curve of each class against the rest, using the uniform class distribution.

Other class probability metrics: [average_precision\(](#page-3-0)), [gain_capture\(](#page-19-0)), [mn_log_loss\(](#page-55-0)), [pr_auc\(](#page-71-0)), [roc_auc\(](#page-81-0)), [roc_aunu\(](#page-88-0))

Examples

```
# Multiclass example
# `obs` is a 4 level factor. The first level is `"VF"`, which is the
# "event of interest" by default in yardstick. See the Relevant Level
# section above.
data(hpc_cv)
# You can use the col1:colN tidyselect syntax
library(dplyr)
hpc_cv %>%
 filter(Resample == "Fold01") %>%
```

```
roc_aunp(obs, VF:L)
# Change the first level of `obs` from `"VF"` to `"M"` to alter the
# event of interest. The class probability columns should be supplied
# in the same order as the levels.
hpc_cv %>%
 filter(Resample == "Fold01") %>%
  mutate(obs = relevel(obs, "M")) %>%
  roc_aunp(obs, M, VF:L)
# Groups are respected
hpc_cv %>%
  group_by(Resample) %>%
  roc_aunp(obs, VF:L)
# Vector version
# Supply a matrix of class probabilities
fold1 <- hpc_cv %>%
  filter(Resample == "Fold01")
roc_aunp_vec(
  truth = fold1$obs,
  matrix(
   c(fold1$VF, fold1$F, fold1$M, fold1$L),
   ncol = 4)
\mathcal{L}# ---------------------------------------------------------------------------
# Options for `pROC::roc()`
# Pass options via a named list and not through `...`!
roc_aunp(
 hpc_cv,
 obs,
 VF:L,
 options = list(smooth = TRUE)
\mathcal{L}
```
roc_aunu *Area under the ROC curve of each class against the rest, using the uniform class distribution*

Description

roc_aunu() is a multiclass metric that computes the area under the ROC curve of each class against the rest, using the uniform class distribution. This is equivalent to roc_auc(estimator $=$ "macro").

Usage

```
roc_aunu(data, ...)
## S3 method for class 'data.frame'
roc_aunu(data, truth, ..., options = list(), na_rm = TRUE)roc_aunu_vec(truth, estimate, options = list(), na_rm = TRUE, ...)
```
Arguments

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For roc_aunu_vec(), a single numeric value (or NA).

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Multiclass

This multiclass method for computing the area under the ROC curve uses the uniform class distribution and is equivalent to roc_auc(estimator = "macro").

Author(s)

Julia Silge

References

Ferri, C., Hernández-Orallo, J., & Modroiu, R. (2009). "An experimental comparison of performance measures for classification". *Pattern Recognition Letters*. 30 (1), pp 27-38.

See Also

[roc_aunp\(\)](#page-85-0) for computing the area under the ROC curve of each class against the rest, using the a priori class distribution.

Other class probability metrics: [average_precision\(](#page-3-0)), [gain_capture\(](#page-19-0)), [mn_log_loss\(](#page-55-0)), [pr_auc\(](#page-71-0)), [roc_auc\(](#page-81-0)), [roc_aunp\(](#page-85-0))

Examples

```
# Multiclass example
```

```
# `obs` is a 4 level factor. The first level is `"VF"`, which is the
# "event of interest" by default in yardstick. See the Relevant Level
# section above.
data(hpc_cv)
# You can use the col1:colN tidyselect syntax
library(dplyr)
hpc_cv %>%
 filter(Resample == "Fold01") %>%
 roc_aunu(obs, VF:L)
# Change the first level of `obs` from `"VF"` to `"M"` to alter the
# event of interest. The class probability columns should be supplied
# in the same order as the levels.
hpc_cv %>%
filter(Resample == "Fold01") %>%
 mutate(obs = relevel(obs, "M")) %>%
```
roc_curve 91

```
roc_aunu(obs, M, VF:L)
# Groups are respected
hpc_cv %>%
  group_by(Resample) %>%
  roc_aunu(obs, VF:L)
# Vector version
# Supply a matrix of class probabilities
fold1 <- hpc_cv %>%
  filter(Resample == "Fold01")
roc_aunu_vec(
  truth = fold1$obs,
  matrix(
    c(fold1$VF, fold1$F, fold1$M, fold1$L),
    ncol = 4)
\overline{\phantom{a}}# ---------------------------------------------------------------------------
# Options for `pROC::roc()`
# Pass options via a named list and not through `...`!
roc_aunu(
  hpc_cv,
  obs,
  VF:L,
  options = list(smooth = TRUE)
\mathcal{L}
```
roc_curve *Receiver operator curve*

Description

roc_curve() constructs the full ROC curve and returns a tibble. See [roc_auc\(\)](#page-81-0) for the area under the ROC curve.

Usage

```
roc_curve(data, ...)
## S3 method for class 'data.frame'
roc_curve(
 data,
 truth,
  ...,
 options = list(),
```

```
na\_rm = TRUE,event_level = yardstick_event_level()
)
```

```
autoplot.roc_df(object, ...)
```
Arguments

Details

roc_curve() computes the sensitivity at every unique value of the probability column (in addition to infinity and minus infinity). If a smooth ROC curve was produced, the unique observed values of the specificity are used to create the curve points. In either case, this may not be efficient for large data sets.

There is a [ggplot2::autoplot\(\)](#page-0-0) method for quickly visualizing the curve. This works for binary and multiclass output, and also works with grouped data (i.e. from resamples). See the examples.

Value

A tibble with class roc_df or roc_grouped_df having columns specificity and sensitivity. If an ordinary (i.e. non-smoothed) curve is used, there is also a column for .threshold.

Multiclass

If a multiclass truth column is provided, a one-vs-all approach will be taken to calculate multiple curves, one per level. In this case, there will be an additional column, .level, identifying the "one" column in the one-vs-all calculation.

roc_curve 93

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Author(s)

Max Kuhn

See Also

Compute the area under the ROC curve with [roc_auc\(\)](#page-81-0).

Other curve metrics: [gain_curve\(](#page-22-0)), [lift_curve\(](#page-38-0)), [pr_curve\(](#page-74-0))

Examples

```
# ---------------------------------------------------------------------------
# Two class example
# 'truth' is a 2 level factor. The first level is '"Class1"', which is the
# "event of interest" by default in yardstick. See the Relevant Level
# section above.
data(two_class_example)
# Binary metrics using class probabilities take a factor `truth` column,
# and a single class probability column containing the probabilities of
# the event of interest. Here, since `"Class1"` is the first level of
# `"truth"`, it is the event of interest and we pass in probabilities for it.
roc_curve(two_class_example, truth, Class1)
# ---------------------------------------------------------------------------
# `autoplot()`
# Visualize the curve using ggplot2 manually
library(ggplot2)
library(dplyr)
roc_curve(two_class_example, truth, Class1) %>%
  ggplot(aes(x = 1 - specificity, y = sensitivity)) +geom_path() +
  geom_abline(lty = 3) +
  coord_equal() +
  theme_bw()
# Or use autoplot
autoplot(roc_curve(two_class_example, truth, Class1))
```
Not run:

```
# Multiclass one-vs-all approach
# One curve per level
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  roc_curve(obs, VF:L) %>%
  autoplot()
# Same as above, but will all of the resamples
hpc_cv %>%
  group_by(Resample) %>%
  roc_curve(obs, VF:L) %>%
  autoplot()
## End(Not run)
```
rpd *Ratio of performance to deviation*

Description

These functions are appropriate for cases where the model outcome is a numeric. The ratio of performance to deviation $(rpd())$ $(rpd())$ $(rpd())$ and the ratio of performance to inter-quartile $(rpiq())$ $(rpiq())$ $(rpiq())$ are both measures of consistency/correlation between observed and predicted values (and not of accuracy).

Usage

```
rpd(data, ...)
## S3 method for class 'data.frame'
rpd(data, truth, estimate, na_rm = TRUE, ...)
```

```
rpd_vec(truth, estimate, na_rm = TRUE, ...)
```
Arguments

Details

In the field of spectroscopy in particular, the ratio of performance to deviation (RPD) has been used as the standard way to report the quality of a model. It is the ratio between the standard deviation of a variable and the standard error of prediction of that variable by a given model. However, its systematic use has been criticized by several authors, since using the standard deviation to represent the spread of a variable can be misleading on skewed dataset. The ratio of performance to interquartile has been introduced by Bellon-Maurel et al. (2010) to address some of these issues, and generalise the RPD to non-normally distributed variables.

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For rpd_vec(), a single numeric value (or NA).

Author(s)

Pierre Roudier

References

Williams, P.C. (1987) Variables affecting near-infrared reflectance spectroscopic analysis. In: Near Infrared Technology in the Agriculture and Food Industries. 1st Ed. P.Williams and K.Norris, Eds. Am. Cereal Assoc. Cereal Chem., St. Paul, MN.

Bellon-Maurel, V., Fernandez-Ahumada, E., Palagos, B., Roger, J.M. and McBratney, A., (2010). Critical review of chemometric indicators commonly used for assessing the quality of the prediction of soil attributes by NIR spectroscopy. TrAC Trends in Analytical Chemistry, 29(9), pp.1073-1081.

See Also

The closely related inter-quartile metric: [rpiq\(\)](#page-95-0)

Other numeric metrics: [ccc\(](#page-9-0)), [huber_loss_pseudo\(](#page-30-0)), [huber_loss\(](#page-28-0)), [iic\(](#page-32-0)), [mae\(](#page-41-0)), [mape\(](#page-43-0)), [mase\(](#page-44-0)), [mpe\(](#page-57-0)), [rmse\(](#page-79-0)), [rpiq\(](#page-95-0)), [rsq_trad\(](#page-99-0)), [rsq\(](#page-97-0)), [smape\(](#page-105-0))

Other consistency metrics: [ccc\(](#page-9-0)), [rpiq\(](#page-95-0)), [rsq_trad\(](#page-99-0)), [rsq\(](#page-97-0))

Examples

Supply truth and predictions as bare column names rpd(solubility_test, solubility, prediction)

```
library(dplyr)
```
set.seed(1234) size <- 100 times <-10

create 10 resamples solubility_resampled <- bind_rows(

rpd 95

```
replicate(
    n = times,
    expr = sample_n(solubility_test, size, replace = TRUE),
    simplify = FALSE
  ),
  .id = "resample"
\mathcal{L}# Compute the metric by group
metric_results <- solubility_resampled %>%
  group_by(resample) %>%
  rpd(solubility, prediction)
metric_results
# Resampled mean estimate
metric_results %>%
  summarise(avg_estimate = mean(.estimate))
```
rpiq *Ratio of performance to inter-quartile*

Description

These functions are appropriate for cases where the model outcome is a numeric. The ratio of performance to deviation ([rpd\(\)](#page-93-0)) and the ratio of performance to inter-quartile ([rpiq\(\)](#page-95-0)) are both measures of consistency/correlation between observed and predicted values (and not of accuracy).

Usage

```
rpiq(data, ...)
## S3 method for class 'data.frame'
rpiq(data, truth, estimate, na_rm = TRUE, ...)
```
rpiq_vec(truth, estimate, na_rm = TRUE, ...)

Arguments

na_rm A logical value indicating whether NA values should be stripped before the computation proceeds.

Details

In the field of spectroscopy in particular, the ratio of performance to deviation (RPD) has been used as the standard way to report the quality of a model. It is the ratio between the standard deviation of a variable and the standard error of prediction of that variable by a given model. However, its systematic use has been criticized by several authors, since using the standard deviation to represent the spread of a variable can be misleading on skewed dataset. The ratio of performance to interquartile has been introduced by Bellon-Maurel et al. (2010) to address some of these issues, and generalise the RPD to non-normally distributed variables.

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For rpd_vec(), a single numeric value (or NA).

Author(s)

Pierre Roudier

References

Williams, P.C. (1987) Variables affecting near-infrared reflectance spectroscopic analysis. In: Near Infrared Technology in the Agriculture and Food Industries. 1st Ed. P.Williams and K.Norris, Eds. Am. Cereal Assoc. Cereal Chem., St. Paul, MN.

Bellon-Maurel, V., Fernandez-Ahumada, E., Palagos, B., Roger, J.M. and McBratney, A., (2010). Critical review of chemometric indicators commonly used for assessing the quality of the prediction of soil attributes by NIR spectroscopy. TrAC Trends in Analytical Chemistry, 29(9), pp.1073-1081.

See Also

The closely related deviation metric: [rpd\(\)](#page-93-0)

Other numeric metrics: [ccc\(](#page-9-0)), [huber_loss_pseudo\(](#page-30-0)), [huber_loss\(](#page-28-0)), [iic\(](#page-32-0)), [mae\(](#page-41-0)), [mape\(](#page-43-0)), [mase\(](#page-44-0)), [mpe\(](#page-57-0)), [rmse\(](#page-79-0)), [rpd\(](#page-93-0)), [rsq_trad\(](#page-99-0)), [rsq\(](#page-97-0)), [smape\(](#page-105-0))

Other consistency metrics: [ccc\(](#page-9-0)), [rpd\(](#page-93-0)), [rsq_trad\(](#page-99-0)), [rsq\(](#page-97-0))

Examples

```
# Supply truth and predictions as bare column names
rpd(solubility_test, solubility, prediction)
```

```
library(dplyr)
```
set.seed(1234) size <- 100 times <-10

```
# create 10 resamples
solubility_resampled <- bind_rows(
  replicate(
    n = times,
    expr = sample_n(solubility_test, size, replace = TRUE),
    simplify = FALSE
  ),
  .id = "resample"
\mathcal{L}# Compute the metric by group
metric_results <- solubility_resampled %>%
  group_by(resample) %>%
  rpd(solubility, prediction)
metric_results
# Resampled mean estimate
metric_results %>%
  summarise(avg_estimate = mean(.estimate))
```
rsq *R squared*

Description

Calculate the coefficient of determination using correlation. For the traditional measure of R squared, see [rsq_trad\(\)](#page-99-0).

Usage

```
rsq(data, ...)
```
S3 method for class 'data.frame' rsq(data, truth, estimate, na_rm = TRUE, ...)

rsq_vec(truth, estimate, na_rm = TRUE, ...)

Arguments

Details

The two estimates for the coefficient of determination, $rsq()$ and $rsq_train()$, differ by their formula. The former guarantees a value on $(0, 1)$ while the latter can generate inaccurate values when the model is non-informative (see the examples). Both are measures of consistency/correlation and not of accuracy.

rsq() is simply the squared correlation between truth and estimate.

Because rsq() internally computes a correlation, if either truth or estimate are constant it can result in a divide by zero error. In these cases, a warning is thrown and NA is returned. This can occur when a model predicts a single value for all samples. For example, a regularized model that eliminates all predictors except for the intercept would do this. Another example would be a CART model that contains no splits.

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For rsq_vec(), a single numeric value (or NA).

Author(s)

Max Kuhn

References

Kvalseth. Cautionary note about R^2 . American Statistician (1985) vol. 39 (4) pp. 279-285.

See Also

Other numeric metrics: [ccc\(](#page-9-0)), [huber_loss_pseudo\(](#page-30-0)), [huber_loss\(](#page-28-0)), [iic\(](#page-32-0)), [mae\(](#page-41-0)), [mape\(](#page-43-0)), [mase\(](#page-44-0)), [mpe\(](#page-57-0)), [rmse\(](#page-79-0)), [rpd\(](#page-93-0)), [rpiq\(](#page-95-0)), [rsq_trad\(](#page-99-0)), [smape\(](#page-105-0))

Other consistency metrics: [ccc\(](#page-9-0)), [rpd\(](#page-93-0)), [rpiq\(](#page-95-0)), [rsq_trad\(](#page-99-0))

Examples

```
# Supply truth and predictions as bare column names
rsq(solubility_test, solubility, prediction)
```
library(dplyr)

set.seed(1234) size <- 100 times <-10

```
# create 10 resamples
solubility_resampled <- bind_rows(
 replicate(
   n = times,
   expr = sample_n(solubility_test, size, replace = TRUE),
   simplify = FALSE
 ),
  .id = "resample"
\lambda# Compute the metric by group
metric_results <- solubility_resampled %>%
 group_by(resample) %>%
 rsq(solubility, prediction)
metric_results
# Resampled mean estimate
metric_results %>%
 summarise(avg_estimate = mean(.estimate))
# With uninformitive data, the traditional version of R^2 can return
# negative values.
set.seed(2291)
solubility_test$randomized <- sample(solubility_test$prediction)
rsq(solubility_test, solubility, randomized)
rsq_trad(solubility_test, solubility, randomized)
# A constant `truth` or `estimate` vector results in a warning from
# a divide by zero error in the correlation calculation.
# `NA` will be returned in these cases.
truth \leq c(1, 2)estimate \leq c(1, 1)
rsq_vec(truth, estimate)
```
rsq_trad *R squared - traditional*

Description

Calculate the coefficient of determination using the traditional definition of R squared using sum of squares. For a measure of R squared that is strictly between $(0, 1)$, see [rsq\(\)](#page-97-0).

Usage

```
rsq_trad(data, ...)
## S3 method for class 'data.frame'
rsq_trad(data, truth, estimate, na_rm = TRUE, ...)
rsq_trad_vec(truth, estimate, na_rm = TRUE, ...)
```
rsq_trad 101

Arguments

Details

The two estimates for the coefficient of determination, $rsq()$ and $rsq_train(d)$, differ by their formula. The former guarantees a value on $(0, 1)$ while the latter can generate inaccurate values when the model is non-informative (see the examples). Both are measures of consistency/correlation and not of accuracy.

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For rsq_trad_vec(), a single numeric value (or NA).

Author(s)

Max Kuhn

References

Kvalseth. Cautionary note about R^2 . American Statistician (1985) vol. 39 (4) pp. 279-285.

See Also

Other numeric metrics: [ccc\(](#page-9-0)), [huber_loss_pseudo\(](#page-30-0)), [huber_loss\(](#page-28-0)), [iic\(](#page-32-0)), [mae\(](#page-41-0)), [mape\(](#page-43-0)), [mase\(](#page-44-0)), [mpe\(](#page-57-0)), [rmse\(](#page-79-0)), [rpd\(](#page-93-0)), [rpiq\(](#page-95-0)), [rsq\(](#page-97-0)), [smape\(](#page-105-0))

Other consistency metrics: [ccc\(](#page-9-0)), [rpd\(](#page-93-0)), [rpiq\(](#page-95-0)), [rsq\(](#page-97-0))

Examples

Supply truth and predictions as bare column names rsq_trad(solubility_test, solubility, prediction)

library(dplyr)

```
set.seed(1234)
size <- 100
times <- 10
# create 10 resamples
solubility_resampled <- bind_rows(
  replicate(
   n = times,
   expr = sample_n(solubility_test, size, replace = TRUE),
   simplify = FALSE
  ),
  .id = "resample"
\mathcal{L}# Compute the metric by group
metric_results <- solubility_resampled %>%
  group_by(resample) %>%
  rsq_trad(solubility, prediction)
metric_results
# Resampled mean estimate
metric_results %>%
  summarise(avg_estimate = mean(.estimate))
# With uninformitive data, the traditional version of R^2 can return
# negative values.
set.seed(2291)
solubility_test$randomized <- sample(solubility_test$prediction)
rsq(solubility_test, solubility, randomized)
rsq_trad(solubility_test, solubility, randomized)
```
sens *Sensitivity*

Description

These functions calculate the [sens\(\)](#page-101-0) (sensitivity) of a measurement system compared to a reference result (the "truth" or gold standard). Highly related functions are $spec(), ppv(),$ $spec(), ppv(),$ $spec(), ppv(),$ $spec(), ppv(),$ and $npv(.)$.

Usage

```
sens(data, ...)
## S3 method for class 'data.frame'
sens(
  data,
  truth,
  estimate,
```

```
estimator = NULL,
  na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
\mathcal{L}sensitivity(data, ...)
sens_vec(
  truth,
  estimate,
  estimator = NULL,
  na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
)
sensitivity_vec(
  truth,
  estimate,
  estimator = NULL,
  na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
\mathcal{L}
```
Arguments

however, if the deprecated global option yardstick.event_first is set, that will be used instead with a warning.

Details

The sensitivity (sens()) is defined as the proportion of positive results out of the number of samples which were actually positive.

When the denominator of the calculation is θ , sensitivity is undefined. This happens when both # true_positive = 0 and # false_negative = 0 are true, which mean that there were no true events. When computing binary sensitivity, a NA value will be returned with a warning. When computing multiclass sensitivity, the individual NA values will be removed, and the computation will procede, with a warning.

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For sens_vec(), a single numeric value (or NA).

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Multiclass

Macro, micro, and macro-weighted averaging is available for this metric. The default is to select macro averaging if a truth factor with more than 2 levels is provided. Otherwise, a standard binary calculation is done. See vignette("multiclass","yardstick") for more information.

Implementation

Suppose a 2x2 table with notation:

The formulas used here are:

Sensitivity =
$$
A/(A+C)
$$

Specificity = $D/(B+D)$

```
sens and \frac{105}{200} 105
```

```
Prevalence = (A+C)/(A+B+C+D)
```

```
P P V = (Sensitivity∗P revalence)/((Sensitivity∗P revalence)+((1−Specif icity)∗(1−P revalence)))
NPV = (Specificity * (1-Prevalence)) / (( (1-Sensitivity) * Prevalence) + ((Specificity) * (1-Prevalence)))
```
See the references for discussions of the statistics.

Author(s)

Max Kuhn

References

Altman, D.G., Bland, J.M. (1994) "Diagnostic tests 1: sensitivity and specificity," *British Medical Journal*, vol 308, 1552.

See Also

```
Other class metrics: accuracy(), bal_accuracy(), detection_prevalence(), f_meas(), j_index(),
kap(), mcc(), npv(), ppv(), precision(), recall(), spec()
```
Other sensitivity metrics: [npv\(](#page-60-0)), [ppv\(](#page-64-0)), [spec\(](#page-107-0))

Examples

```
# Two class
data("two_class_example")
sens(two_class_example, truth, predicted)
# Multiclass
library(dplyr)
data(hpc_cv)
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  sens(obs, pred)
# Groups are respected
hpc_cv %>%
  group_by(Resample) %>%
  sens(obs, pred)
# Weighted macro averaging
hpc_cv %>%
  group_by(Resample) %>%
  sens(obs, pred, estimator = "macro_weighted")
# Vector version
sens_vec(
  two_class_example$truth,
  two_class_example$predicted
)
```
106 smape smape

```
# Making Class2 the "relevant" level
sens_vec(
  two_class_example$truth,
  two_class_example$predicted,
  event_level = "second"
\mathcal{L}
```


smape *Symmetric mean absolute percentage error*

Description

Calculate the symmetric mean absolute percentage error. This metric is in *relative units*.

Usage

```
smape(data, ...)
## S3 method for class 'data.frame'
smape(data, truth, estimate, na_rm = TRUE, ...)
```

```
smape_vec(truth, estimate, na_rm = TRUE, ...)
```
Arguments

Details

This implementation of smape() is the "usual definition" where the denominator is divided by two.

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For smape_vec(), a single numeric value (or NA).

solubility_test 107

Author(s)

Max Kuhn, Riaz Hedayati

See Also

```
Other numeric metrics: ccc(), huber_loss_pseudo(), huber_loss(), iic(), mae(), mape(),
mase(), mpe(), rmse(), rpd(), rpiq(), rsq_trad(), rsq()
```

```
Other accuracy metrics: ccc(), huber_loss_pseudo(), huber_loss(), iic(), mae(), mape(),
mase(), mpe(), rmse()
```
Examples

```
# Supply truth and predictions as bare column names
smape(solubility_test, solubility, prediction)
library(dplyr)
set.seed(1234)
size <- 100
times <- 10
# create 10 resamples
solubility_resampled <- bind_rows(
  replicate(
   n = times,
   expr = sample_n(solubility_test, size, replace = TRUE),
   simplify = FALSE
  ),
  .id = "resample"
)
# Compute the metric by group
metric_results <- solubility_resampled %>%
  group_by(resample) %>%
  smape(solubility, prediction)
metric_results
# Resampled mean estimate
metric_results %>%
  summarise(avg_estimate = mean(.estimate))
```
solubility_test *Solubility Predictions from MARS Model*

Description

Solubility Predictions from MARS Model

Details

For the solubility data in Kuhn and Johnson (2013), these data are the test set results for the MARS model. The observed solubility (in column solubility) and the model results (prediction) are contained in the data.

Value

solubility_test

a data frame

Source

Kuhn, M., Johnson, K. (2013) *Applied Predictive Modeling*, Springer

Examples

data(solubility_test) str(solubility_test)

spec *Specificity*

Description

These functions calculate the [spec\(\)](#page-107-0) (specificity) of a measurement system compared to a reference result (the "truth" or gold standard). Highly related functions are [sens\(\)](#page-101-0), [ppv\(\)](#page-64-0), and [npv\(\)](#page-60-0).

Usage

```
spec(data, ...)
## S3 method for class 'data.frame'
spec(
 data,
 truth,
 estimate,
 estimator = NULL,
 na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
\lambdaspecificity(data, ...)
spec_vec(
  truth,
 estimate,
```
```
\sec 109
```

```
estimator = NULL,
 na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
\mathcal{L}specificity_vec(
  truth,
 estimate,
 estimator = NULL,
 na\_rm = TRUE,event_level = yardstick_event_level(),
  ...
)
```
Arguments

Details

The specificity measures the proportion of negatives that are correctly identified as negatives.

When the denominator of the calculation is 0 , specificity is undefined. This happens when both # true_negative $= 0$ and $\#$ false_positive $= 0$ are true, which mean that there were no true negatives. When computing binary specificity, a NA value will be returned with a warning. When computing multiclass specificity, the individual NA values will be removed, and the computation will procede, with a warning.

Value

A tibble with columns .metric, .estimator, and .estimate and 1 row of values.

For grouped data frames, the number of rows returned will be the same as the number of groups.

For spec_vec(), a single numeric value (or NA).

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

Multiclass

Macro, micro, and macro-weighted averaging is available for this metric. The default is to select macro averaging if a truth factor with more than 2 levels is provided. Otherwise, a standard binary calculation is done. See vignette("multiclass","yardstick") for more information.

Implementation

Suppose a 2x2 table with notation:

The formulas used here are:

 $Sensitivity = A/(A+C)$

$$
Specificity = D/(B+D)
$$

 $Prevalence = (A+C)/(A+B+C+D)$

 $PPV = (Sensitivity*Prevalence)/((Sensitivity*Prevalence)+((1-Specificity)*(1-Prevalence)))$

 $NPV = (Specificity * (1-Prevalence)) / (((1-Sensitivity) * Prevalence) + ((Specificity) * (1-Prevalence)))$

See the references for discussions of the statistics.

Author(s)

Max Kuhn

References

Altman, D.G., Bland, J.M. (1994) "Diagnostic tests 1: sensitivity and specificity," *British Medical Journal*, vol 308, 1552.

See Also

Other class metrics: [accuracy\(](#page-2-0)), [bal_accuracy\(](#page-6-0)), [detection_prevalence\(](#page-13-0)), [f_meas\(](#page-16-0)), [j_index\(](#page-34-0)), [kap\(](#page-37-0)), [mcc\(](#page-47-0)), [npv\(](#page-60-0)), [ppv\(](#page-64-0)), [precision\(](#page-67-0)), [recall\(](#page-76-0)), [sens\(](#page-101-0))

Other sensitivity metrics: [npv\(](#page-60-0)), [ppv\(](#page-64-0)), [sens\(](#page-101-0))

Examples

```
# Two class
data("two_class_example")
spec(two_class_example, truth, predicted)
# Multiclass
library(dplyr)
data(hpc_cv)
hpc_cv %>%
  filter(Resample == "Fold01") %>%
  spec(obs, pred)
# Groups are respected
hpc_cv %>%
  group_by(Resample) %>%
  spec(obs, pred)
# Weighted macro averaging
hpc_cv %>%
  group_by(Resample) %>%
  spec(obs, pred, estimator = "macro_weighted")
# Vector version
spec_vec(
  two_class_example$truth,
  two_class_example$predicted
)
# Making Class2 the "relevant" level
spec_vec(
  two_class_example$truth,
  two_class_example$predicted,
  event_level = "second"
\lambda
```


Description

Various statistical summaries of confusion matrices are produced and returned in a tibble. These include those shown in the help pages for [sens\(\)](#page-101-0), [recall\(\)](#page-76-0), and [accuracy\(\)](#page-2-0), among others.

Usage

```
## S3 method for class 'conf_mat'
summary(
 object,
 prevalence = NULL,
 beta = 1,
 estimator = NULL,
 event_level = yardstick_event_level(),
  ...
)
```
Arguments

Value

A tibble containing various classification metrics.

Relevant Level

There is no common convention on which factor level should automatically be considered the "event" or "positive" result when computing binary classification metrics. In yardstick, the default is to use the *first* level. To alter this, change the argument event_level to "second" to consider

two_class_example 113

the *last* level of the factor the level of interest. For multiclass extensions involving one-vs-all comparisons (such as macro averaging), this option is ignored and the "one" level is always the relevant result.

See Also

[conf_mat\(\)](#page-11-0)

Examples

```
data("two_class_example")
cmat <- conf_mat(two_class_example, truth = "truth", estimate = "predicted")
summary(cmat)
summary(cmat, prevalence = 0.70)
library(dplyr)
library(purrr)
library(tidyr)
data("hpc_cv")
# Compute statistics per resample then summarize
all_metrics <- hpc_cv %>%
 group_by(Resample) %>%
 conf_mat(obs, pred) %>%
 mutate(summary_tbl = map(conf_mat, summary)) %>%
 unnest(summary_tbl)
all_metrics %>%
 group_by(.metric) %>%
 summarise(
   mean = mean(.estimate, na.rm = TRUE),
   sd = sd(.estimate, na.rm = TRUE)
 )
```
two_class_example *Two Class Predictions*

Description

Two Class Predictions

Details

These data are a test set form a model built for two classes ("Class1" and "Class2"). There are columns for the true and predicted classes and column for the probabilities for each class.

Value

two_class_example

a data frame

Examples

```
data(two_class_example)
str(two_class_example)
```
'truth' is a 2 level factor. The first level is '"Class1"', which is the # "event of interest" by default in yardstick. See the Relevant Level # section in any classification function (such as `?pr_auc`) to see how # to change this. levels(hpc_cv\$obs)

Index

∗ accuracy metrics ccc , [10](#page-9-0) huber_loss , [29](#page-28-0) huber_loss_pseudo , [31](#page-30-0) iic , [33](#page-32-0) mae , [42](#page-41-0) mape , [44](#page-43-0) mase , [45](#page-44-0) mpe , [58](#page-57-0) rmse , [80](#page-79-0) smape , [106](#page-105-0) ∗ class metrics accuracy , [3](#page-2-1) bal_accuracy , [7](#page-6-1) detection_prevalence , [14](#page-13-1) f_meas , [17](#page-16-1) j_index , [35](#page-34-1) kap , [38](#page-37-1) mcc , [48](#page-47-1) npv , [61](#page-60-1) ppv , [65](#page-64-1) precision , [68](#page-67-1) recall , [77](#page-76-1) sens , [102](#page-101-1) spec , [108](#page-107-0) ∗ class probability metrics average_precision , [4](#page-3-0) gain_capture, $20\,$ $20\,$ mn_log_loss, [56](#page-55-0) pr_auc , [72](#page-71-0) roc_auc , [82](#page-81-0) roc_aunp , [86](#page-85-0) roc_aunu , [89](#page-88-0) ∗ consistency metrics ccc , [10](#page-9-0) rpd , [94](#page-93-0) rpiq , [96](#page-95-0) rsq , [98](#page-97-0) rsq_trad , [100](#page-99-0)

∗ curve metrics gain_curve , [23](#page-22-0) lift_curve , [39](#page-38-0) pr_curve , [75](#page-74-0) roc_curve , [91](#page-90-0) ∗ datasets hpc_cv, [28](#page-27-0) pathology , [64](#page-63-0) solubility_test , [107](#page-106-0) two_class_example , [113](#page-112-0) ∗ numeric metrics ccc , [10](#page-9-0) huber_loss , [29](#page-28-0) huber_loss_pseudo , [31](#page-30-0) iic , [33](#page-32-0) mae , [42](#page-41-0) mape , [44](#page-43-0) mase , [45](#page-44-0) mpe , [58](#page-57-0) rmse , [80](#page-79-0) rpd , [94](#page-93-0) rpiq , [96](#page-95-0) rsq , [98](#page-97-0) rsq_trad , [100](#page-99-0) smape, [106](#page-105-0) ∗ relevance metrics f_meas , [17](#page-16-1) precision , [68](#page-67-1) recall , [77](#page-76-1) ∗ sensitivity metrics npv , [61](#page-60-1) ppv , [65](#page-64-1) sens , [102](#page-101-1) spec, [108](#page-107-0) accuracy , [3](#page-2-1) , *[9](#page-8-0)* , *[16](#page-15-0)* , *[19](#page-18-0)* , *[37](#page-36-0)* , *[39](#page-38-0)* , *[49](#page-48-0)* , *[63](#page-62-0)* , *[67](#page-66-0)* , *[71](#page-70-0)* , *[80](#page-79-0)* , *[105](#page-104-0)* , *[111](#page-110-0)* accuracy() , *[38](#page-37-1)* , *[50](#page-49-0)* , *[57](#page-56-0)* , *[112](#page-111-0)* accuracy_vec *(*accuracy *)* , [3](#page-2-1)

autoplot.conf_mat *(*conf_mat *)* , [12](#page-11-1)

116 INDEX

autoplot.gain_df *(*gain_curve*)*, [23](#page-22-0) autoplot.lift_df *(*lift_curve*)*, [39](#page-38-0) autoplot.pr_df *(*pr_curve*)*, [75](#page-74-0) autoplot.roc_df *(*roc_curve*)*, [91](#page-90-0) average_precision, [4,](#page-3-0) *[22](#page-21-0)*, *[58](#page-57-0)*, *[73](#page-72-0)*, *[84](#page-83-0)*, *[87](#page-86-0)*, *[90](#page-89-0)* average_precision_vec *(*average_precision*)*, [4](#page-3-0)

bal_accuracy, *[4](#page-3-0)*, [7,](#page-6-1) *[16](#page-15-0)*, *[19](#page-18-0)*, *[37](#page-36-0)*, *[39](#page-38-0)*, *[49](#page-48-0)*, *[63](#page-62-0)*, *[67](#page-66-0)*, *[71](#page-70-0)*, *[80](#page-79-0)*, *[105](#page-104-0)*, *[111](#page-110-0)* bal_accuracy_vec *(*bal_accuracy*)*, [7](#page-6-1) base::table(), *[12](#page-11-1)*

ccc, [10,](#page-9-0) *[30](#page-29-0)*, *[32](#page-31-0)*, *[34](#page-33-0)*, *[43](#page-42-0)*, *[45](#page-44-0)*, *[47](#page-46-0)*, *[59](#page-58-0)*, *[81](#page-80-0)*, *[95](#page-94-0)*, *[97](#page-96-0)*, *[99](#page-98-0)*, *[101](#page-100-0)*, *[107](#page-106-0)* ccc(), *[11](#page-10-0)* ccc_vec *(*ccc*)*, [10](#page-9-0) conf_mat, [12](#page-11-1) conf_mat(), *[13](#page-12-0)*, *[112,](#page-111-0) [113](#page-112-0)*

detection_prevalence, *[4](#page-3-0)*, *[9](#page-8-0)*, [14,](#page-13-1) *[19](#page-18-0)*, *[37](#page-36-0)*, *[39](#page-38-0)*, *[49](#page-48-0)*, *[63](#page-62-0)*, *[67](#page-66-0)*, *[71](#page-70-0)*, *[80](#page-79-0)*, *[105](#page-104-0)*, *[111](#page-110-0)* detection_prevalence_vec *(*detection_prevalence*)*, [14](#page-13-1) developer-helpers *(*get_weights*)*, [26](#page-25-0) dots_to_estimate *(*get_weights*)*, [26](#page-25-0) dots_to_estimate(), *[54–](#page-53-0)[56](#page-55-0)*

f_meas, *[4](#page-3-0)*, *[9](#page-8-0)*, *[16](#page-15-0)*, [17,](#page-16-1) *[37](#page-36-0)*, *[39](#page-38-0)*, *[49](#page-48-0)*, *[63](#page-62-0)*, *[67](#page-66-0)*, *[71](#page-70-0)*, *[80](#page-79-0)*, *[105](#page-104-0)*, *[111](#page-110-0)* f_meas(), *[17](#page-16-1)*, *[68](#page-67-1)*, *[77](#page-76-1)*, *[112](#page-111-0)* f_meas_vec *(*f_meas*)*, [17](#page-16-1) finalize_estimator *(*get_weights*)*, [26](#page-25-0) finalize_estimator(), *[55,](#page-54-0) [56](#page-55-0)* finalize_estimator_internal *(*get_weights*)*, [26](#page-25-0)

gain_capture, *[6](#page-5-0)*, [20,](#page-19-0) *[58](#page-57-0)*, *[73](#page-72-0)*, *[84](#page-83-0)*, *[87](#page-86-0)*, *[90](#page-89-0)* gain_capture(), *[23](#page-22-0)*, *[25](#page-24-0)* gain_capture_vec *(*gain_capture*)*, [20](#page-19-0) gain_curve, [23,](#page-22-0) *[41](#page-40-0)*, *[76](#page-75-0)*, *[93](#page-92-0)* gain_curve(), *[22](#page-21-0)*, *[39](#page-38-0)* get_weights, [26](#page-25-0) ggplot2::autoplot(), *[13](#page-12-0)*, *[24](#page-23-0)*, *[40](#page-39-0)*, *[76](#page-75-0)*, *[92](#page-91-0)*

hpc_cv, 28 huber_loss, *[11](#page-10-0)*, [29,](#page-28-0) *[32](#page-31-0)*, *[34](#page-33-0)*, *[43](#page-42-0)*, *[45](#page-44-0)*, *[47](#page-46-0)*, *[59](#page-58-0)*, *[81](#page-80-0)*, *[95](#page-94-0)*, *[97](#page-96-0)*, *[99](#page-98-0)*, *[101](#page-100-0)*, *[107](#page-106-0)* huber_loss(), *[31](#page-30-0)*

huber_loss_pseudo, *[11](#page-10-0)*, *[30](#page-29-0)*, [31,](#page-30-0) *[34](#page-33-0)*, *[43](#page-42-0)*, *[45](#page-44-0)*, *[47](#page-46-0)*, *[59](#page-58-0)*, *[81](#page-80-0)*, *[95](#page-94-0)*, *[97](#page-96-0)*, *[99](#page-98-0)*, *[101](#page-100-0)*, *[107](#page-106-0)* huber_loss_pseudo_vec *(*huber_loss_pseudo*)*, [31](#page-30-0) huber_loss_vec *(*huber_loss*)*, [29](#page-28-0) iic, *[11](#page-10-0)*, *[30](#page-29-0)*, *[32](#page-31-0)*, [33,](#page-32-0) *[43](#page-42-0)*, *[45](#page-44-0)*, *[47](#page-46-0)*, *[59](#page-58-0)*, *[81](#page-80-0)*, *[95](#page-94-0)*, *[97](#page-96-0)*, *[99](#page-98-0)*, *[101](#page-100-0)*, *[107](#page-106-0)* iic_vec *(*iic*)*, [33](#page-32-0) j_index, *[4](#page-3-0)*, *[9](#page-8-0)*, *[16](#page-15-0)*, *[19](#page-18-0)*, [35,](#page-34-1) *[39](#page-38-0)*, *[49](#page-48-0)*, *[63](#page-62-0)*, *[67](#page-66-0)*, *[71](#page-70-0)*, *[80](#page-79-0)*, *[105](#page-104-0)*, *[111](#page-110-0)* j_index_vec *(*j_index*)*, [35](#page-34-1) kap, *[4](#page-3-0)*, *[9](#page-8-0)*, *[16](#page-15-0)*, *[19](#page-18-0)*, *[37](#page-36-0)*, [38,](#page-37-1) *[49](#page-48-0)*, *[63](#page-62-0)*, *[67](#page-66-0)*, *[71](#page-70-0)*, *[80](#page-79-0)*, *[105](#page-104-0)*, *[111](#page-110-0)* kap(), *[50](#page-49-0)* kap_vec *(*kap*)*, [38](#page-37-1) lift_curve, *[25](#page-24-0)*, [39,](#page-38-0) *[76](#page-75-0)*, *[93](#page-92-0)* lift_curve(), *[23](#page-22-0)* mae, *[11](#page-10-0)*, *[30](#page-29-0)*, *[32](#page-31-0)*, *[34](#page-33-0)*, [42,](#page-41-0) *[45](#page-44-0)*, *[47](#page-46-0)*, *[59](#page-58-0)*, *[81](#page-80-0)*, *[95](#page-94-0)*, *[97](#page-96-0)*, *[99](#page-98-0)*, *[101](#page-100-0)*, *[107](#page-106-0)* mae(), *[50](#page-49-0)* mae_vec *(*mae*)*, [42](#page-41-0) mape, *[11](#page-10-0)*, *[30](#page-29-0)*, *[32](#page-31-0)*, *[34](#page-33-0)*, *[43](#page-42-0)*, [44,](#page-43-0) *[47](#page-46-0)*, *[59](#page-58-0)*, *[81](#page-80-0)*, *[95](#page-94-0)*, *[97](#page-96-0)*, *[99](#page-98-0)*, *[101](#page-100-0)*, *[107](#page-106-0)* mape_vec *(*mape*)*, [44](#page-43-0) mase, *[11](#page-10-0)*, *[30](#page-29-0)*, *[32](#page-31-0)*, *[34](#page-33-0)*, *[43](#page-42-0)*, *[45](#page-44-0)*, [45,](#page-44-0) *[59](#page-58-0)*, *[81](#page-80-0)*, *[95](#page-94-0)*, *[97](#page-96-0)*, *[99](#page-98-0)*, *[101](#page-100-0)*, *[107](#page-106-0)* mase_vec *(*mase*)*, [45](#page-44-0) mcc, *[4](#page-3-0)*, *[9](#page-8-0)*, *[16](#page-15-0)*, *[19](#page-18-0)*, *[37](#page-36-0)*, *[39](#page-38-0)*, [48,](#page-47-1) *[63](#page-62-0)*, *[67](#page-66-0)*, *[71](#page-70-0)*, *[80](#page-79-0)*, *[105](#page-104-0)*, *[111](#page-110-0)* mcc_vec *(*mcc*)*, [48](#page-47-1) metric_set, [51](#page-50-0) metric_set(), *[50](#page-49-0)*, *[60](#page-59-0)* metric_summarizer, [53](#page-52-0) metric_summarizer(), *[26](#page-25-0)*, *[28](#page-27-0)*, *[55,](#page-54-0) [56](#page-55-0)* metric_vec_template, [55](#page-54-0) metric_vec_template(), *[26](#page-25-0)*, *[28](#page-27-0)*, *[53](#page-52-0)*, *[55](#page-54-0)* metrics, [49](#page-48-0) metrics(), *[52](#page-51-0)* mn_log_loss, *[6](#page-5-0)*, *[22](#page-21-0)*, [56,](#page-55-0) *[73](#page-72-0)*, *[84](#page-83-0)*, *[87](#page-86-0)*, *[90](#page-89-0)* mn_log_loss(), *[50](#page-49-0)* mn_log_loss_vec *(*mn_log_loss*)*, [56](#page-55-0) mpe, *[11](#page-10-0)*, *[30](#page-29-0)*, *[32](#page-31-0)*, *[34](#page-33-0)*, *[43](#page-42-0)*, *[45](#page-44-0)*, *[47](#page-46-0)*, [58,](#page-57-0) *[81](#page-80-0)*, *[95](#page-94-0)*, *[97](#page-96-0)*, *[99](#page-98-0)*, *[101](#page-100-0)*, *[107](#page-106-0)* mpe_vec *(*mpe*)*, [58](#page-57-0)

INDEX 117

new-metric, [60](#page-59-0) new_class_metric *(*new-metric*)*, [60](#page-59-0) new_numeric_metric *(*new-metric*)*, [60](#page-59-0) new_prob_metric *(*new-metric*)*, [60](#page-59-0) npv, *[4](#page-3-0)*, *[9](#page-8-0)*, *[16](#page-15-0)*, *[19](#page-18-0)*, *[37](#page-36-0)*, *[39](#page-38-0)*, *[49](#page-48-0)*, [61,](#page-60-1) *[67](#page-66-0)*, *[71](#page-70-0)*, *[80](#page-79-0)*, *[105](#page-104-0)*, *[111](#page-110-0)* npv(), *[61,](#page-60-1) [62](#page-61-0)*, *[65,](#page-64-1) [66](#page-65-0)*, *[102](#page-101-1)*, *[108](#page-107-0)* npv_vec *(*npv*)*, [61](#page-60-1)

pathology, [64](#page-63-0) ppv, *[4](#page-3-0)*, *[9](#page-8-0)*, *[16](#page-15-0)*, *[19](#page-18-0)*, *[37](#page-36-0)*, *[39](#page-38-0)*, *[49](#page-48-0)*, *[63](#page-62-0)*, [65,](#page-64-1) *[71](#page-70-0)*, *[80](#page-79-0)*, *[105](#page-104-0)*, *[111](#page-110-0)* ppv(), *[61,](#page-60-1) [62](#page-61-0)*, *[65,](#page-64-1) [66](#page-65-0)*, *[102](#page-101-1)*, *[108](#page-107-0)* ppv_vec *(*ppv*)*, [65](#page-64-1) pr_auc, *[6](#page-5-0)*, *[22](#page-21-0)*, *[58](#page-57-0)*, [72,](#page-71-0) *[84](#page-83-0)*, *[87](#page-86-0)*, *[90](#page-89-0)* pr_auc(), *[6](#page-5-0)*, *[75,](#page-74-0) [76](#page-75-0)* pr_auc_vec *(*pr_auc*)*, [72](#page-71-0) pr_curve, *[25](#page-24-0)*, *[41](#page-40-0)*, [75,](#page-74-0) *[93](#page-92-0)* pr_curve(), *[4](#page-3-0)[–6](#page-5-0)*, *[72,](#page-71-0) [73](#page-72-0)* precision, *[4](#page-3-0)*, *[9](#page-8-0)*, *[16](#page-15-0)*, *[19](#page-18-0)*, *[37](#page-36-0)*, *[39](#page-38-0)*, *[49](#page-48-0)*, *[63](#page-62-0)*, *[67](#page-66-0)*, [68,](#page-67-1) *[80](#page-79-0)*, *[105](#page-104-0)*, *[111](#page-110-0)* precision(), *[17](#page-16-1)*, *[68](#page-67-1)*, *[77](#page-76-1)* precision_vec *(*precision*)*, [68](#page-67-1) pROC::roc(), *[50](#page-49-0)*, *[83](#page-82-0)*, *[87](#page-86-0)*, *[89](#page-88-0)*, *[92](#page-91-0)*

quasiquotation, *[3](#page-2-1)*, *[5](#page-4-0)*, *[8](#page-7-0)*, *[10](#page-9-0)*, *[12](#page-11-1)*, *[15](#page-14-0)*, *[17](#page-16-1)*, *[21](#page-20-0)*, , *[29](#page-28-0)*, *[31](#page-30-0)*, *[33](#page-32-0)*, *[35](#page-34-1)*, *[38](#page-37-1)*, *[40](#page-39-0)*, *[43,](#page-42-0) [44](#page-43-0)*, *[46](#page-45-0)*, , *[50](#page-49-0)*, *[57](#page-56-0)*, *[59](#page-58-0)*, *[62](#page-61-0)*, *[66](#page-65-0)*, *[69](#page-68-0)*, *[72](#page-71-0)*, *[75](#page-74-0)*, *[78](#page-77-0)*, , *[83](#page-82-0)*, *[86](#page-85-0)*, *[89](#page-88-0)*, *[92](#page-91-0)*, *[94](#page-93-0)*, *[96](#page-95-0)*, *[98](#page-97-0)*, *[101](#page-100-0)*, , *[106](#page-105-0)*, *[109](#page-108-0)*

recall, *[4](#page-3-0)*, *[9](#page-8-0)*, *[16](#page-15-0)*, *[19](#page-18-0)*, *[37](#page-36-0)*, *[39](#page-38-0)*, *[49](#page-48-0)*, *[63](#page-62-0)*, *[67](#page-66-0)*, *[71](#page-70-0)*, [77,](#page-76-1) *[105](#page-104-0)*, *[111](#page-110-0)* recall(), *[17](#page-16-1)*, *[68](#page-67-1)*, *[77](#page-76-1)*, *[112](#page-111-0)* recall_vec *(*recall*)*, [77](#page-76-1) rmse, *[11](#page-10-0)*, *[30](#page-29-0)*, *[32](#page-31-0)*, *[34](#page-33-0)*, *[43](#page-42-0)*, *[45](#page-44-0)*, *[47](#page-46-0)*, *[59](#page-58-0)*, [80,](#page-79-0) *[95](#page-94-0)*, *[97](#page-96-0)*, *[99](#page-98-0)*, *[101](#page-100-0)*, *[107](#page-106-0)* rmse(), *[11](#page-10-0)*, *[29](#page-28-0)*, *[31](#page-30-0)*, *[50](#page-49-0)* rmse_vec *(*rmse*)*, [80](#page-79-0) roc_auc, *[6](#page-5-0)*, *[22](#page-21-0)*, *[58](#page-57-0)*, *[73](#page-72-0)*, [82,](#page-81-0) *[87](#page-86-0)*, *[90](#page-89-0)* roc_auc(), *[50](#page-49-0)*, *[91](#page-90-0)*, *[93](#page-92-0)* roc_auc_vec *(*roc_auc*)*, [82](#page-81-0) roc_aunp, *[6](#page-5-0)*, *[22](#page-21-0)*, *[58](#page-57-0)*, *[73](#page-72-0)*, *[84](#page-83-0)*, [86,](#page-85-0) *[90](#page-89-0)* roc_aunp(), *[90](#page-89-0)* roc_aunp_vec *(*roc_aunp*)*, [86](#page-85-0) roc_aunu, *[6](#page-5-0)*, *[22](#page-21-0)*, *[58](#page-57-0)*, *[73](#page-72-0)*, *[84](#page-83-0)*, *[87](#page-86-0)*, [89](#page-88-0) roc_aunu(), *[87](#page-86-0)* roc_aunu_vec *(*roc_aunu*)*, [89](#page-88-0) roc_curve, *[25](#page-24-0)*, *[41](#page-40-0)*, *[76](#page-75-0)*, [91](#page-90-0)

roc_curve(), *[82](#page-81-0)*, *[84](#page-83-0)* rpd, *[11](#page-10-0)*, *[30](#page-29-0)*, *[32](#page-31-0)*, *[34](#page-33-0)*, *[43](#page-42-0)*, *[45](#page-44-0)*, *[47](#page-46-0)*, *[59](#page-58-0)*, *[81](#page-80-0)*, [94,](#page-93-0) *[97](#page-96-0)*, *[99](#page-98-0)*, *[101](#page-100-0)*, *[107](#page-106-0)* rpd(), *[94](#page-93-0)*, *[96,](#page-95-0) [97](#page-96-0)* rpd_vec *(*rpd*)*, [94](#page-93-0) rpiq, *[11](#page-10-0)*, *[30](#page-29-0)*, *[32](#page-31-0)*, *[34](#page-33-0)*, *[43](#page-42-0)*, *[45](#page-44-0)*, *[47](#page-46-0)*, *[59](#page-58-0)*, *[81](#page-80-0)*, *[95](#page-94-0)*, [96,](#page-95-0) *[99](#page-98-0)*, *[101](#page-100-0)*, *[107](#page-106-0)* rpiq(), *[94](#page-93-0)[–96](#page-95-0)* rpiq_vec *(*rpiq*)*, [96](#page-95-0) rsq, *[11](#page-10-0)*, *[30](#page-29-0)*, *[32](#page-31-0)*, *[34](#page-33-0)*, *[43](#page-42-0)*, *[45](#page-44-0)*, *[47](#page-46-0)*, *[59](#page-58-0)*, *[81](#page-80-0)*, *[95](#page-94-0)*, *[97](#page-96-0)*, [98,](#page-97-0) *[101](#page-100-0)*, *[107](#page-106-0)* rsq(), *[11](#page-10-0)*, *[50](#page-49-0)*, *[99](#page-98-0)[–101](#page-100-0)* rsq_trad, *[11](#page-10-0)*, *[30](#page-29-0)*, *[32](#page-31-0)*, *[34](#page-33-0)*, *[43](#page-42-0)*, *[45](#page-44-0)*, *[47](#page-46-0)*, *[59](#page-58-0)*, *[81](#page-80-0)*, *[95](#page-94-0)*, *[97](#page-96-0)*, *[99](#page-98-0)*, [100,](#page-99-0) *[107](#page-106-0)* rsq_trad(), *[98,](#page-97-0) [99](#page-98-0)*, *[101](#page-100-0)* rsq_trad_vec *(*rsq_trad*)*, [100](#page-99-0) rsq_vec *(*rsq*)*, [98](#page-97-0) sens, *[4](#page-3-0)*, *[9](#page-8-0)*, *[16](#page-15-0)*, *[19](#page-18-0)*, *[37](#page-36-0)*, *[39](#page-38-0)*, *[49](#page-48-0)*, *[63](#page-62-0)*, *[67](#page-66-0)*, *[71](#page-70-0)*, *[80](#page-79-0)*, [102,](#page-101-1) *[111](#page-110-0)* sens(), *[8](#page-7-0)*, *[35](#page-34-1)*, *[61](#page-60-1)*, *[65](#page-64-1)*, *[102](#page-101-1)*, *[108](#page-107-0)*, *[112](#page-111-0)* sens_vec *(*sens*)*, [102](#page-101-1) sensitivity *(*sens*)*, [102](#page-101-1) sensitivity_vec *(*sens*)*, [102](#page-101-1) smape, *[11](#page-10-0)*, *[30](#page-29-0)*, *[32](#page-31-0)*, *[34](#page-33-0)*, *[43](#page-42-0)*, *[45](#page-44-0)*, *[47](#page-46-0)*, *[59](#page-58-0)*, *[81](#page-80-0)*, *[95](#page-94-0)*, *[97](#page-96-0)*, *[99](#page-98-0)*, *[101](#page-100-0)*, [106](#page-105-0) smape_vec *(*smape*)*, [106](#page-105-0) solubility_test, [107](#page-106-0) spec, *[4](#page-3-0)*, *[9](#page-8-0)*, *[16](#page-15-0)*, *[19](#page-18-0)*, *[37](#page-36-0)*, *[39](#page-38-0)*, *[49](#page-48-0)*, *[63](#page-62-0)*, *[67](#page-66-0)*, *[71](#page-70-0)*, *[80](#page-79-0)*, *[105](#page-104-0)*, [108](#page-107-0) spec(), *[8](#page-7-0)*, *[35](#page-34-1)*, *[61](#page-60-1)*, *[65](#page-64-1)*, *[102](#page-101-1)*, *[108](#page-107-0)* spec_vec *(*spec*)*, [108](#page-107-0) specificity *(*spec*)*, [108](#page-107-0) specificity_vec *(*spec*)*, [108](#page-107-0) summary.conf_mat, [112](#page-111-0) summary.conf_mat(), *[13](#page-12-0)*

tidy.conf_mat *(*conf_mat*)*, [12](#page-11-1) two_class_example, [113](#page-112-0)

validate_estimator *(*get_weights*)*, [26](#page-25-0)