

Package: SLmetrics (via r-universe)

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Title Machine Learning Performance Evaluation on Steroids

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Description Performance evaluation metrics for supervised and unsupervised machine learning, statistical learning and artificial intelligence applications. Core computations are implemented in 'C++' for scalability and efficiency.

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<https://github.com/serkor1/SLmetrics>

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| | |
|-----------------|-----------------|
| accuracy.factor | <i>Accuracy</i> |
|-----------------|-----------------|

Description

A generic function for the (normalized) **accuracy** in classification tasks. Use `weighted.accuracy()` for the weighted **accuracy**.

Usage

```
## S3 method for class 'factor'
accuracy(actual, predicted, ...)

## S3 method for class 'factor'
weighted.accuracy(actual, predicted, w, ...)

## S3 method for class 'cmatrix'
accuracy(x, ...)

## Generic S3 method
accuracy(...)

## Generic S3 method
weighted.accuracy(
  ...,
  w
)
```

Arguments

| | |
|-----------|--|
| actual | A vector of <code><factor></code> with length n , and k levels |
| predicted | A vector of <code><factor></code> with length n , and k levels |
| ... | Arguments passed into other methods |
| w | A <code><numeric></code> -vector of length n . <code>NULL</code> by default |
| x | A confusion matrix created <code>cmatrix()</code> |

Value

A `<numeric>`-vector of **length** 1

Definition

Let $\hat{\alpha} \in [0, 1]$ be the proportion of correctly predicted classes. The **accuracy** of the classifier is calculated as,

$$\hat{\alpha} = \frac{\#TP + \#TN}{\#TP + \#TN + \#FP + \#FN}$$

Where:

- $\#TP$ is the number of true positives,
- $\#TN$ is the number of true negatives,
- $\#FP$ is the number of false positives, and
- $\#FN$ is the number of false negatives.

Creating <factor>

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

See Also

Other Classification: `ROC.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zeroone.loss.factor()`

Other Supervised Learning: `ROC.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroone.loss.factor()`

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data = iris,
  family = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
predicted <- factor(
  as.numeric(
    predict(model, type = "response") > 0.5
  ),
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)
```

```
# 4) evaluate model
# performance
cat(
  "Accuracy", accuracy(
    actual = actual,
    predicted = predicted
  ),
  "Accuracy (weigthed)", weighted.accuracy(
    actual = actual,
    predicted = predicted,
    w      = iris$Petal.Length/mean(iris$Petal.Length)
  ),
  sep = "\n"
)
```

 auc.numeric

 AUC

Description

The `auc()`-function calculates the area under the curve.

Usage

```
## S3 method for class 'numeric'
auc(y, x, method = 0L, presorted = TRUE, ...)

## Generic S3 method
auc(
  y,
  x,
  method = 0,
  presorted = TRUE,
  ...
)
```

Arguments

| | |
|------------------------|--|
| <code>y</code> | A <numeric> vector of length <i>n</i> . |
| <code>x</code> | A <numeric> vector of length <i>n</i> . |
| <code>method</code> | A <numeric> value (default: 0). Defines the underlying method of calculating the area under the curve. If 0 it is calculated using the trapezoid-method, if 1 it is calculated using the step-method. |
| <code>presorted</code> | A <logical> -value length 1 (default: FALSE). If TRUE the input will not be sorted by threshold. |
| <code>...</code> | Arguments passed into other methods. |

Value

A <numeric> vector of length 1

Definition**Trapezoidal rule**

The **trapezoidal rule** approximates the integral of a function $f(x)$ between $x = a$ and $x = b$ using trapezoids formed between consecutive points. If we have points x_0, x_1, \dots, x_n (with $a = x_0 < x_1 < \dots < x_n = b$) and corresponding function values $f(x_0), f(x_1), \dots, f(x_n)$, the area under the curve A_T is approximated by:

$$A_T \approx \sum_{k=1}^n \frac{f(x_{k-1}) + f(x_k)}{2} [x_k - x_{k-1}].$$

Step-function method

The **step-function (rectangular) method** uses the value of the function at one endpoint of each subinterval to form rectangles. With the same partition x_0, x_1, \dots, x_n , the rectangular approximation A_S can be written as:

$$A_S \approx \sum_{k=1}^n f(x_{k-1}) [x_k - x_{k-1}].$$

See Also

Other Tools: [cov.wt.matrix\(\)](#), [preorder\(\)](#), [presort\(\)](#)

Examples

```
## 1) Ordered x and y pair
x <- seq(0, pi, length.out = 200)
y <- sin(x)

## 1.1) calculate area
ordered_auc <- auc(y = y, x = x)

## 2) Unordered x and y pair
x <- sample(seq(0, pi, length.out = 200))
y <- sin(x)

## 2.1) calculate area
unordered_auc <- auc(y = y, x = x)

## 2.2) calculate area with explicit
## ordering
unordered_auc_flag <- auc(
  y = y,
  x = x,
  presorted = FALSE
)
```

```
## 3) display result
cat(
  "AUC (ordered x and y pair)", ordered_auc,
  "AUC (unordered x and y pair)", unordered_auc,
  "AUC (unordered x and y pair, with unordered flag)", unordered_auc_flag,
  sep = "\n"
)
```

| | |
|------------------|--------------------------|
| baccuracy.factor | <i>Balanced Accuracy</i> |
|------------------|--------------------------|

Description

A generic function for the (normalized) balanced **accuracy**. Use `weighted.baccuracy()` for the weighted balanced **accuracy**.

Usage

```
## S3 method for class 'factor'
baccuracy(actual, predicted, adjust = FALSE, na.rm = TRUE, ...)

## S3 method for class 'factor'
weighted.baccuracy(actual, predicted, w, adjust = FALSE, na.rm = TRUE, ...)

## S3 method for class 'cmatrix'
baccuracy(x, adjust = FALSE, na.rm = TRUE, ...)

## Generic S3 method
baccuracy(
  ...,
  adjust = FALSE,
  na.rm = TRUE
)

## Generic S3 method
weighted.baccuracy(
  ...,
  w,
  adjust = FALSE,
  na.rm = TRUE
)
```

Arguments

| | |
|-----------|---|
| actual | A vector of <code><factor></code> with length n , and k levels |
| predicted | A vector of <code><factor></code> with length n , and k levels |

| | |
|--------|--|
| adjust | A logical value (default: FALSE). If TRUE the metric is adjusted for random chance $\frac{1}{k}$. |
| na.rm | A logical value (default: TRUE). If TRUE calculation of the metric is based on valid classes. |
| ... | Arguments passed into other methods |
| w | A <numeric> -vector of length n . NULL by default |
| x | A confusion matrix created <code>cmatrix()</code> |

Value

A **numeric**-vector of length 1

Definition

Let $\hat{\alpha} \in [0, 1]$ be the proportion of correctly predicted classes. If `adjust == false`, the balanced **accuracy** of the classifier is calculated as,

$$\hat{\alpha} = \frac{\text{sensitivity} + \text{specificity}}{2}$$

otherwise,

$$\hat{\alpha} = \frac{\text{sensitivity} + \text{specificity}}{2} \frac{1}{k}$$

Where:

- k is the number of classes
- sensitivity is the overall **sensitivity**, and
- specificity is the overall **specificity**

Creating <factor>

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

See Also

Other Classification: `ROC.factor()`, `accuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zeroone.loss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroone.loss.factor()`

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data = iris,
  family = binomial(
    link = "logit"
  )
)
```

```
)

# 3) generate predicted
# classes
predicted <- factor(
  as.numeric(
    predict(model, type = "response") > 0.5
  ),
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 4) evaluate the
# model
cat(
  "Balanced accuracy", baccuracy(
    actual = actual,
    predicted = predicted
  ),
  "Balanced accuracy (weigthed)", weighted.baccuracy(
    actual = actual,
    predicted = predicted,
    w = iris$Petal.Length/mean(iris$Petal.Length)
  ),
  sep = "\n"
)
```

banknote

Banknote Authentication Dataset

Description

This dataset contains features extracted from the wavelet transform of banknote images, which are used to classify banknotes as authentic or inauthentic. The data originates from the UCI Machine Learning Repository.

Usage

`data(banknote)`

Format

A list with two components:

features A data frame with 4 variables: variance, skewness, curtosis, and entropy.

target A factor with levels "inauthentic" and "authentic" representing the banknote's authenticity.

Details

The data is provided as a list with two components:

features A data frame containing the following variables:

variance Variance of the wavelet transformed image.

skewness Skewness of the wavelet transformed image.

curtosis Curtosis of the wavelet transformed image.

entropy Entropy of the image.

target A factor indicating the authenticity of the banknote. The factor has two levels:

inauthentic Indicates the banknote is not genuine.

authentic Indicates the banknote is genuine.

Source

<https://archive.ics.uci.edu/dataset/267/banknote+authentication>

ccc.numeric

Concordance Correlation Coefficient

Description

A generic function for the **concordance correlation coefficient**. Use `weighted.ccc()` for the weighted **concordance correlation coefficient**.

Usage

```
## S3 method for class 'numeric'
ccc(actual, predicted, correction = FALSE, ...)

## S3 method for class 'numeric'
weighted.ccc(actual, predicted, w, correction = FALSE, ...)

ccc(
  ...,
  correction = FALSE
)

weighted.ccc(
```

```

    ...,
    w,
    correction = FALSE
)

```

Arguments

actual A <numeric>-vector of length n . The observed (continuous) response variable.

predicted A <numeric>-vector of length n . The estimated (continuous) response variable.

correction A <logical> vector of length 1 (default: `FALSE`). If `TRUE` the variance and covariance will be adjusted with $\frac{1-n}{n}$.

... Arguments passed into other methods.

w A <numeric>-vector of length n . The weight assigned to each observation in the data.

Value

A <numeric> vector of length 1.

Definition

Let $\rho_c \in [0, 1]$ measure the agreement between y and v . The classifier agreement is calculated as,

$$\rho_c = \frac{2\rho\sigma_v\sigma_y}{\sigma_v^2 + \sigma_y^2 + (\mu_v - \mu_y)^2}$$

Where:

- ρ is the pearson correlation coefficient
- σ_y is the unbiased standard deviation of y
- σ_v is the unbiased standard deviation of v
- μ_y is the mean of y
- μ_v is the mean of v

If correction == TRUE each $\sigma_{i \in \{y,v\}}$ is adjusted by $\frac{1-n}{n}$

See Also

Other Regression: `huberloss.numeric()`, `mae.numeric()`, `mape.numeric()`, `mpe.numeric()`, `mse.numeric()`, `pinball.numeric()`, `rae.numeric()`, `rmse.numeric()`, `rmsle.numeric()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroone.loss.factor()`

Examples

```

# 1) fit a linear
# regression
model <- lm(
  mpg ~ .,
  data = mtcars
)

# 1.1) define actual
# and predicted values
# to measure performance
actual <- mtcars$mpg
predicted <- fitted(model)

# 2) evaluate in-sample model
# performance
cat(
  "Concordance Correlation Coefficient", ccc(
    actual = actual,
    predicted = predicted,
    correction = FALSE
  ),
  "Concordance Correlation Coefficient (corrected)", ccc(
    actual = actual,
    predicted = predicted,
    correction = TRUE
  ),
  "Concordance Correlation Coefficient (weighed)", weighted.ccc(
    actual = actual,
    predicted = predicted,
    w = mtcars$mpg/mean(mtcars$mpg),
    correction = FALSE
  ),
  sep = "\n"
)

```

ckappa.factor

Cohen's κ -statistic

Description

A generic function for **Cohen's κ -statistic**. Use `weighted.ckappa()` for the weighted κ -statistic.

Usage

```

## S3 method for class 'factor'
ckappa(actual, predicted, beta = 0, ...)

## S3 method for class 'factor'

```

```

weighted.ckappa(actual, predicted, w, beta = 0, ...)

## S3 method for class 'cmatrix'
ckappa(x, beta = 0, ...)

ckappa(
  ...,
  beta = 0
)

weighted.ckappa(
  ...,
  w,
  beta = 0
)

```

Arguments

| | |
|-----------|--|
| actual | A vector of <factor> values of length n , and k levels. |
| predicted | A vector of <factor> values of length n , and k levels. |
| beta | A <numeric> value of length 1 (default: 0). If $\beta \neq 0$ the off-diagonals of the confusion matrix are penalized with a factor of $(y_+ - y_{i,-})^\beta$. |
| ... | Arguments passed into other methods |
| w | A <numeric>-vector of length n . NULL by default. |
| x | A confusion matrix created <code>cmatrix()</code> . |

Value

If `micro` is `NULL` (the default), a named <numeric>-vector of length k

If `micro` is `TRUE` or `FALSE`, a <numeric>-vector of length 1

Definition

Let $\kappa \in [0, 1]$ be the inter-rater (intra-rater) reliability. The inter-rater (intra-rater) reliability is calculated as,

$$\kappa = \frac{\rho_p - \rho_e}{1 - \rho_e}$$

Where:

- ρ_p is the empirical probability of agreement between predicted and actual values
- ρ_e is the expected probability of agreement under random chance

If $\beta \neq 0$ the off-diagonals in the confusion matrix is penalized before ρ is calculated. More formally,

$$\chi = X \circ Y^\beta$$

Where:

- X is the confusion matrix
- Y is the penalizing matrix and
- β is the penalizing factor

Creating <factor>

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

See Also

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zeroonloss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`,


```
mape.numeric(), mcc.factor(), mpe.numeric(), mse.numeric(), nlr.factor(), npv.factor(),  
pinball.numeric(), plr.factor(), pr.auc.matrix(), prROC.factor(), precision.factor(),  
rae.numeric(), recall.factor(), rmse.numeric(), rmsle.numeric(), roc.auc.matrix(),  
rrmse.numeric(), rrse.numeric(), rsq.numeric(), smape.numeric(), specificity.factor(),  
zeroone.loss.factor()
```

Examples

```
# 1) recode Iris  
# to binary classification  
# problem  
iris$species_num <- as.numeric(  
  iris$Species == "virginica"  
)  
  
# 2) fit the logistic  
# regression  
model <- glm(  
  formula = species_num ~ Sepal.Length + Sepal.Width,  
  data = iris,  
  family = binomial(  
    link = "logit"  
  )  
)  
  
# 3) generate predicted  
# classes  
predicted <- factor(  
  as.numeric(  
    predict(model, type = "response") > 0.5  
  ),  
  levels = c(1,0),  
  labels = c("Virginica", "Others")  
)  
  
# 3.1) generate actual  
# classes  
actual <- factor(  
  x = iris$species_num,  
  levels = c(1,0),  
  labels = c("Virginica", "Others")  
)  
  
# 4) evaluate model performance with  
# Cohens Kappa statistic  
cat(  
  "Kappa", ckappa(  
    actual = actual,  
    predicted = predicted  
  ),  
  "Kappa (penalized)", ckappa(  
    actual = actual,
```

```

    predicted = predicted,
    beta      = 2
  ),
  "Kappa (weigthed)", weighted.ckappa(
    actual    = actual,
    predicted = predicted,
    w         = iris$Petal.Length/mean(iris$Petal.Length)
  ),
  sep = "\n"
)

```

cmatrix.factor

Confusion Matrix

Description

The `cmatrix()`-function uses cross-classifying factors to build a confusion matrix of the counts at each combination of the `factor` levels. Each row of the `matrix` represents the actual `factor` levels, while each column represents the predicted `factor` levels.

Usage

```

## S3 method for class 'factor'
cmatrix(actual, predicted, ...)

## S3 method for class 'factor'
weighted.cmatrix(actual, predicted, w, ...)

## Generic S3 method
cmatrix(
  actual,
  predicted,
  ...
)

## Generic S3 method
weighted.cmatrix(
  actual,
  predicted,
  w,
  ...
)

```

Arguments

`actual` A `<factor>`-vector of `length` n , and k levels.
`predicted` A `<factor>`-vector of `length` n , and k levels.

... Arguments passed into other methods.
 w A <numeric>-vector of length n (default: `NULL`) If passed it will return a weighted confusion matrix.

Value

A named $k \times k$ <matrix>

Dimensions

There is no robust defensive measure against mis-specifying the confusion matrix. If the arguments are correctly specified, the resulting confusion matrix is on the form:

| | A (Predicted) | B (Predicted) |
|------------|---------------|---------------|
| A (Actual) | Value | Value |
| B (Actual) | Value | Value |

Creating <factor>

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

See Also

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zeroone.loss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroone.loss.factor()`

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data = iris,
  family = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
predicted <- factor(
  as.numeric(
    predict(model, type = "response") > 0.5
  ),
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)
```

```
# 4) summarise performance
# in a confusion matrix

# 4.1) unweighted matrix
confusion_matrix <- cmatrix(
  actual    = actual,
  predicted = predicted
)

# 4.1.1) summarise matrix
summary(
  confusion_matrix
)

# 4.1.2) plot confusion
# matrix
plot(
  confusion_matrix
)

# 4.2) weighted matrix
confusion_matrix <- weighted.cmatrix(
  actual    = actual,
  predicted = predicted,
  w         = iris$Petal.Length/mean(iris$Petal.Length)
)

# 4.2.1) summarise matrix
summary(
  confusion_matrix
)

# 4.2.1) plot confusion
# matrix
plot(
  confusion_matrix
)
```

dor.factor

Diagnostic Odds Ratio

Description

A generic function for the **diagnostic odds ratio** in classification tasks. Use `weighted.dor()` weighted **diagnostic odds ratio**.

Usage

```
## S3 method for class 'factor'
```

```

dor(actual, predicted, ...)

## S3 method for class 'factor'
weighted.dor(actual, predicted, w, ...)

## S3 method for class 'cmatrix'
dor(x, ...)

## Generic S3 method
dor(...)

## Generic S3 method
weighted.dor(
  ...,
  w
)

```

Arguments

| | |
|-----------|---|
| actual | A vector of <factor> values of length n , and k levels. |
| predicted | A vector of <factor> values of length n , and k levels. |
| ... | Arguments passed into other methods |
| w | A <numeric>-vector of length n . NULL by default. |
| x | A confusion matrix created <code>cmatrix()</code> . |

Value

A <numeric>-vector of length 1

Definition

Let $\hat{\alpha} \in [0, \infty]$ be the effectiveness of the classifier. The **diagnostic odds ratio** of the classifier is calculated as,

$$\hat{\alpha} = \frac{\#TP\#TN}{\#FP\#FN}$$

Where:

- #TP is the number of true positives
- #TN is the number of true negatives
- #FP is the number of false positives
- #FN is the number of false negatives

Creating <factor>

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

See Also

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zeroone.loss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroone.loss.factor()`

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data    = iris,
  family  = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
predicted <- factor(
  as.numeric(
    predict(model, type = "response") > 0.5
  ),
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 4) evaluate model performance
# with Diagnostic Odds Ratio
cat("Diagnostic Odds Ratio", sep = "\n")
dor(
  actual    = actual,
  predicted = predicted
)

cat("Diagnostic Odds Ratio (weighted)", sep = "\n")
weighted.dor(
  actual    = actual,
  predicted = predicted,
  w         = iris$Petal.Length/mean(iris$Petal.Length)
)
```

| | |
|----------------|----------------|
| entropy.matrix | <i>Entropy</i> |
|----------------|----------------|

Description

The `entropy()` function calculates the **Entropy** of given probability distributions.

Usage

```
## S3 method for class 'matrix'
entropy(pk, dim = 0L, base = -1, ...)

## S3 method for class 'matrix'
relative.entropy(pk, qk, dim = 0L, base = -1, ...)

## S3 method for class 'matrix'
cross.entropy(pk, qk, dim = 0L, base = -1, ...)

## Generic S3 method
entropy(
  pk,
  dim = 0,
  base = -1,
  ...
)

## Generic S3 method
relative.entropy(
  pk,
  qk,
  dim = 0,
  base = -1,
  ...
)

## Generic S3 method
cross.entropy(
  pk,
  qk,
  dim = 0,
  base = -1,
  ...
)
```

Arguments

`pk` A $n \times k$ **<numeric>**-matrix of observed probabilities. The i -th row should sum to 1 (i.e., a valid probability distribution over the k classes). The first column

| | |
|------|---|
| | corresponds to the first factor level in actual, the second column to the second factor level, and so on. |
| dim | An <code><integer></code> value of <code>length</code> 1 (Default: 0). Defines the dimension along which to calculate the entropy (0: total, 1: row-wise, 2: column-wise). |
| base | A <code><numeric></code> value of <code>length</code> 1 (Default: -1). The logarithmic base to use. Default value specifies natural logarithms. |
| ... | Arguments passed into other methods |
| qk | A $n \times k$ <code><numeric></code> -matrix of predicted probabilities. The i -th row should sum to 1 (i.e., a valid probability distribution over the k classes). The first column corresponds to the first factor level in actual, the second column to the second factor level, and so on. |

Value

A `<numeric>` value or vector:

- A single `<numeric>` value (length 1) if `dim == 0`.
- A `<numeric>` vector with length equal to the `length` of rows if `dim == 1`.
- A `<numeric>` vector with length equal to the `length` of columns if `dim == 2`.

Definition

Entropy:

$$H(pk) = - \sum_i pk_i \log(pk_i)$$

Cross Entropy:

$$H(pk, qk) = - \sum_i pk_i \log(qk_i)$$

Relative Entropy

$$D_{KL}(pk \parallel qk) = \sum_i pk_i \log \left(\frac{pk_i}{qk_i} \right)$$

See Also

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zeroone.loss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroone.loss.factor()`

Examples

```

# 1) Define actual
# and observed probabilities

# 1.1) actual probabilities
pk <- matrix(
  cbind(1/2, 1/2),
  ncol = 2
)

# 1.2) observed (estimated) probabilities
qk <- matrix(
  cbind(9/10, 1/10),
  ncol = 2
)

# 2) calculate
# Entropy
cat(
  "Entropy", entropy(
    pk
  ),
  "Relative Entropy", relative.entropy(
    pk,
    qk
  ),
  "Cross Entropy", cross.entropy(
    pk,
    qk
  ),
  sep = "\n"
)

```

fbeta.factor

 F_{β} -score

Description

A generic function for the F_{β} -score. Use `weighted.fbeta()` for the weighted F_{β} -score.

Usage

```

## S3 method for class 'factor'
fbeta(actual, predicted, beta = 1, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
weighted.fbeta(actual, predicted, w, beta = 1, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'cmatrix'

```

```
fbeta(x, beta = 1, micro = NULL, na.rm = TRUE, ...)

## Generic S3 method
fbeta(
  ...,
  beta = 1,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
weighted.fbeta(
  ...,
  w,
  beta = 1,
  micro = NULL,
  na.rm = TRUE
)
```

Arguments

| | |
|-----------|---|
| actual | A vector of <factor> values of length n , and k levels. |
| predicted | A vector of <factor> values of length n , and k levels. |
| beta | A <numeric> vector of length 1 (default: 1). |
| micro | A <logical>-value of length 1 (default: <code>NULL</code>). If <code>TRUE</code> it returns the micro average across all k classes, if <code>FALSE</code> it returns the macro average. |
| na.rm | A <logical> value of length 1 (default: <code>TRUE</code>). If <code>TRUE</code> , <code>NA</code> values are removed from the computation. This argument is only relevant when <code>micro != NULL</code> . When <code>na.rm = TRUE</code> , the computation corresponds to <code>sum(c(1, 2, NA), na.rm = TRUE) / length(na.omit(c(1, 2, NA)))</code> . When <code>na.rm = FALSE</code> , the computation corresponds to <code>sum(c(1, 2, NA), na.rm = TRUE) / length(c(1, 2, NA))</code> . |
| ... | Arguments passed into other methods |
| w | A <numeric>-vector of length n . <code>NULL</code> by default. |
| x | A confusion matrix created <code>cmatrix()</code> . |

Value

If `micro` is `NULL` (the default), a named <numeric>-vector of length k

If `micro` is `TRUE` or `FALSE`, a <numeric>-vector of length 1

Definition

Let $\hat{F}_\beta \in [0, 1]$ be the F_β score, which is a weighted harmonic mean of precision and recall. F_β score of the classifier is calculated as,

$$\hat{F}_\beta = (1 + \beta^2) \frac{\text{Precision} \times \text{Recall}}{\beta^2 \times \text{Precision} + \text{Recall}}$$

Substituting Precision = $\frac{\#TP_k}{\#TP_k + \#FP_k}$ and Recall = $\frac{\#TP_k}{\#TP_k + \#FN_k}$ yields:

$$\hat{F}_\beta = (1 + \beta^2) \frac{\frac{\#TP_k}{\#TP_k + \#FP_k} \times \frac{\#TP_k}{\#TP_k + \#FN_k}}{\beta^2 \times \frac{\#TP_k}{\#TP_k + \#FP_k} + \frac{\#TP_k}{\#TP_k + \#FN_k}}$$

Where:

- $\#TP_k$ is the number of true positives,
- $\#FP_k$ is the number of false positives,
- $\#FN_k$ is the number of false negatives, and
- β is a non-negative real number that determines the relative importance of precision vs. recall in the score.

Creating `<factor>`

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

See Also

Other Classification: [ROC.factor\(\)](#), [accuracy.factor\(\)](#), [baccuracy.factor\(\)](#), [ckappa.factor\(\)](#), [cmatrix.factor\(\)](#), [dor.factor\(\)](#), [entropy.matrix\(\)](#), [fdr.factor\(\)](#), [fer.factor\(\)](#), [fmi.factor\(\)](#), [fpr.factor\(\)](#), [jaccard.factor\(\)](#), [logloss.factor\(\)](#), [mcc.factor\(\)](#), [nlr.factor\(\)](#), [npv.factor\(\)](#), [plr.factor\(\)](#), [pr.auc.matrix\(\)](#), [prROC.factor\(\)](#), [precision.factor\(\)](#), [recall.factor\(\)](#), [roc.auc.matrix\(\)](#), [specificity.factor\(\)](#), [zeroone.loss.factor\(\)](#)

Other Supervised Learning: [ROC.factor\(\)](#), [accuracy.factor\(\)](#), [baccuracy.factor\(\)](#), [ccc.numeric\(\)](#), [ckappa.factor\(\)](#), [cmatrix.factor\(\)](#), [dor.factor\(\)](#), [entropy.matrix\(\)](#), [fdr.factor\(\)](#), [fer.factor\(\)](#), [fpr.factor\(\)](#), [huberloss.numeric\(\)](#), [jaccard.factor\(\)](#), [logloss.factor\(\)](#), [mae.numeric\(\)](#), [mape.numeric\(\)](#), [mcc.factor\(\)](#), [mpe.numeric\(\)](#), [mse.numeric\(\)](#), [nlr.factor\(\)](#), [npv.factor\(\)](#), [pinball.numeric\(\)](#), [plr.factor\(\)](#), [pr.auc.matrix\(\)](#), [prROC.factor\(\)](#), [precision.factor\(\)](#), [rae.numeric\(\)](#), [recall.factor\(\)](#), [rmse.numeric\(\)](#), [rmsle.numeric\(\)](#), [roc.auc.matrix\(\)](#), [rrmse.numeric\(\)](#), [rrse.numeric\(\)](#), [rsq.numeric\(\)](#), [smape.numeric\(\)](#), [specificity.factor\(\)](#), [zeroone.loss.factor\(\)](#)

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data = iris,
  family = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
predicted <- factor(
  as.numeric(
    predict(model, type = "response") > 0.5
  ),
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)
```

```

# 4) evaluate class-wise performance
# using F1-score

# 4.1) unweighted F1-score
fbeta(
  actual    = actual,
  predicted  = predicted,
  beta      = 1
)

# 4.2) weighted F1-score
weighted.fbeta(
  actual    = actual,
  predicted  = predicted,
  w         = iris$Petal.Length/mean(iris$Petal.Length),
  beta      = 1
)

# 5) evaluate overall performance
# using micro-averaged F1-score
cat(
  "Micro-averaged F1-score", fbeta(
    actual    = actual,
    predicted  = predicted,
    beta      = 1,
    micro     = TRUE
  ),
  "Micro-averaged F1-score (weighted)", weighted.fbeta(
    actual    = actual,
    predicted  = predicted,
    w         = iris$Petal.Length/mean(iris$Petal.Length),
    beta      = 1,
    micro     = TRUE
  ),
  sep = "\n"
)

```

| | |
|------------|-----------------------------|
| fdr.factor | <i>false discovery rate</i> |
|------------|-----------------------------|

Description

A generic function for the **False Discovery Rate**. Use `weighted.fdr()` for the weighted **False Discovery Rate**.

Usage

```

## S3 method for class 'factor'
fdr(actual, predicted, micro = NULL, na.rm = TRUE, ...)

```

```

## S3 method for class 'factor'
weighted.fdr(actual, predicted, w, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'cmatrix'
fdr(x, micro = NULL, na.rm = TRUE, ...)

## Generic S3 method
fdr(
  ...,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
weighted.fdr(
  ...,
  w,
  micro = NULL,
  na.rm = TRUE
)

```

Arguments

| | |
|-----------|--|
| actual | A vector of <code><factor></code> values of length n , and k levels. |
| predicted | A vector of <code><factor></code> values of length n , and k levels. |
| micro | A <code><logical></code> -value of length 1 (default: <code>NULL</code>). If <code>TRUE</code> it returns the micro average across all k classes, if <code>FALSE</code> it returns the macro average. |
| na.rm | A <code><logical></code> value of length 1 (default: <code>TRUE</code>). If <code>TRUE</code> , <code>NA</code> values are removed from the computation. This argument is only relevant when <code>micro != NULL</code> . When <code>na.rm = TRUE</code> , the computation corresponds to <code>sum(c(1, 2, NA), na.rm = TRUE) / length(na.omit(c(1, 2, NA)))</code> . When <code>na.rm = FALSE</code> , the computation corresponds to <code>sum(c(1, 2, NA), na.rm = TRUE) / length(c(1, 2, NA))</code> . |
| ... | Arguments passed into other methods |
| w | A <code><numeric></code> -vector of length n . <code>NULL</code> by default. |
| x | A confusion matrix created <code>cmatrix()</code> . |

Value

If `micro` is `NULL` (the default), a named `<numeric>`-vector of length k
 If `micro` is `TRUE` or `FALSE`, a `<numeric>`-vector of length 1

Definition

Let $\hat{\alpha} \in [0, 1]$ be the proportion of false positives among the predicted positives. The false discovery rate of the classifier is calculated as,

$$\hat{\alpha} = \frac{\#FP_k}{\#TP_k + \#FP_k}$$

Where:

- $\#TP_k$ is the number of true positives, and
- $\#FP_k$ is the number of false positives

Creating `<factor>`

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

See Also

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zeroonloss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroone.loss.factor()`

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data = iris,
  family = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
predicted <- factor(
  as.numeric(
    predict(model, type = "response") > 0.5
  ),
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 4) evaluate class-wise performance
# using False Discovery Rate

# 4.1) unweighted False Discovery Rate
fdr(
  actual = actual,
```

```

    predicted = predicted
  )

# 4.2) weighted False Discovery Rate
weighted.fdr(
  actual    = actual,
  predicted = predicted,
  w         = iris$Petal.Length/mean(iris$Petal.Length)
)

# 5) evaluate overall performance
# using micro-averaged False Discovery Rate
cat(
  "Micro-averaged False Discovery Rate", fdr(
    actual    = actual,
    predicted = predicted,
    micro     = TRUE
  ),
  "Micro-averaged False Discovery Rate (weighted)", weighted.fdr(
    actual    = actual,
    predicted = predicted,
    w         = iris$Petal.Length/mean(iris$Petal.Length),
    micro     = TRUE
  ),
  sep = "\n"
)

```

fer.factor

False Omission Rate

Description

A generic function for the **false omission rate**. Use `weighted.fdr()` for the weighted **false omission rate**.

Usage

```

## S3 method for class 'factor'
fer(actual, predicted, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
weighted.fer(actual, predicted, w, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'cmatrix'
fer(x, micro = NULL, na.rm = TRUE, ...)

## Generic S3 method
fer(

```

```

    ...,
    micro = NULL,
    na.rm = TRUE
  )

## Generic S3 method
weighted.fer(
  ...,
  w,
  micro = NULL,
  na.rm = TRUE
)

```

Arguments

| | |
|-----------|--|
| actual | A vector of <code><factor></code> values of length n , and k levels. |
| predicted | A vector of <code><factor></code> values of length n , and k levels. |
| micro | A <code><logical></code> -value of length 1 (default: <code>NULL</code>). If <code>TRUE</code> it returns the micro average across all k classes, if <code>FALSE</code> it returns the macro average. |
| na.rm | A <code><logical></code> value of length 1 (default: <code>TRUE</code>). If <code>TRUE</code> , <code>NA</code> values are removed from the computation. This argument is only relevant when <code>micro != NULL</code> . When <code>na.rm = TRUE</code> , the computation corresponds to <code>sum(c(1, 2, NA), na.rm = TRUE) / length(na.omit(c(1, 2, NA)))</code> . When <code>na.rm = FALSE</code> , the computation corresponds to <code>sum(c(1, 2, NA), na.rm = TRUE) / length(c(1, 2, NA))</code> . |
| ... | Arguments passed into other methods |
| w | A <code><numeric></code> -vector of length n . <code>NULL</code> by default. |
| x | A confusion matrix created <code>cmatrix()</code> . |

Value

If `micro` is `NULL` (the default), a named `<numeric>`-vector of length k

If `micro` is `TRUE` or `FALSE`, a `<numeric>`-vector of length 1

Definition

Let $\hat{\beta} \in [0, 1]$ be the proportion of false negatives among the predicted negatives. The false omission rate of the classifier is calculated as,

$$\hat{\beta} = \frac{\#FN_k}{\#TN_k + \#FN_k}$$

Where:

- $\#TN_k$ is the number of true negatives, and
- $\#FN_k$ is the number of false negatives.

Creating <factor>

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

See Also

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zeroone.loss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroone.loss.factor()`

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data    = iris,
  family  = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
predicted <- factor(
  as.numeric(
    predict(model, type = "response") > 0.5
  ),
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 4) evaluate class-wise performance
# using False Omission Rate

# 4.1) unweighted False Omission Rate
fer(
  actual    = actual,
  predicted = predicted
)

# 4.2) weighted False Omission Rate
weighted.fer(
  actual    = actual,
  predicted = predicted,
  w         = iris$Petal.Length/mean(iris$Petal.Length)
)
```

```

# 5) evaluate overall performance
# using micro-averaged False Omission Rate
cat(
  "Micro-averaged False Omission Rate", fer(
    actual = actual,
    predicted = predicted,
    micro = TRUE
  ),
  "Micro-averaged False Omission Rate (weighted)", weighted.fer(
    actual = actual,
    predicted = predicted,
    w = iris$Petal.Length/mean(iris$Petal.Length),
    micro = TRUE
  ),
  sep = "\n"
)

```

fmi.factor

Fowlkes-Mallows Index

Description

The `fmi()`-function computes the **Fowlkes-Mallows Index** (FMI), a measure of the similarity between two sets of clusterings, between two vectors of predicted and observed `factor()` values.

Usage

```

## S3 method for class 'factor'
fmi(actual, predicted, ...)

## S3 method for class 'cmatrix'
fmi(x, ...)

## Generic S3 method
fmi(...)

```

Arguments

| | |
|-----------|---|
| actual | A vector of <code><factor></code> with <code>length</code> n , and k levels |
| predicted | A vector of <code><factor></code> with <code>length</code> n , and k levels |
| ... | Arguments passed into other methods |
| x | A confusion matrix created <code>cmatrix()</code> |

Value

A `<numeric>`-vector of `length` 1

Definition

The metric is calculated for each class k as follows,

$$\sqrt{\frac{\#TP_k}{\#TP_k + \#FP_k} \times \frac{\#TP_k}{\#TP_k + \#FN_k}}$$

Where $\#TP_k$, $\#FP_k$, and $\#FN_k$ represent the number of true positives, false positives, and false negatives for each class k , respectively.

Creating <factor>

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

See Also

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zeroonloss.factor()`

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data = iris,
  family = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
predicted <- factor(
  as.numeric(
    predict(model, type = "response") > 0.5
  ),
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 4) evaluate model performance
# using Fowlkes Mallows Index
cat(
  "Fowlkes Mallows Index", fmi(
    actual = actual,
    predicted = predicted
  ),
  sep = "\n"
)
```

Description

A generic function for the **False Positive Rate**. Use `weighted.fpr()` for the weighted **False Positive Rate**.

Other names:

Fallout

Usage

```
## S3 method for class 'factor'
fpr(actual, predicted, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
weighted.fpr(actual, predicted, w, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'cmatrix'
fpr(x, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
fallout(actual, predicted, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
weighted.fallout(actual, predicted, w, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'cmatrix'
fallout(x, micro = NULL, na.rm = TRUE, ...)

## Generic S3 method
fpr(
  ...,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
fallout(
  ...,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
weighted.fpr(
  ...,
  w,
  micro = NULL,
  na.rm = TRUE
)
```

```
## Generic S3 method
weighted.fallout(
  ...,
  w,
  micro = NULL,
  na.rm = TRUE
)
```

Arguments

| | |
|-----------|---|
| actual | A vector of <factor> values of length n , and k levels. |
| predicted | A vector of <factor> values of length n , and k levels. |
| micro | A <logical>-value of length 1 (default: <code>NULL</code>). If <code>TRUE</code> it returns the micro average across all k classes, if <code>FALSE</code> it returns the macro average. |
| na.rm | A <logical> value of length 1 (default: <code>TRUE</code>). If <code>TRUE</code> , <code>NA</code> values are removed from the computation. This argument is only relevant when <code>micro != NULL</code> . When <code>na.rm = TRUE</code> , the computation corresponds to <code>sum(c(1, 2, NA), na.rm = TRUE) / length(na.omit(c(1, 2, NA)))</code> . When <code>na.rm = FALSE</code> , the computation corresponds to <code>sum(c(1, 2, NA), na.rm = TRUE) / length(c(1, 2, NA))</code> . |
| ... | Arguments passed into other methods |
| w | A <numeric>-vector of length n . <code>NULL</code> by default. |
| x | A confusion matrix created <code>cmatrix()</code> . |

Value

If `micro` is `NULL` (the default), a named <numeric>-vector of length k

If `micro` is `TRUE` or `FALSE`, a <numeric>-vector of length 1

Definition

Let $\hat{\gamma} \in [0, 1]$ be the proportion of false positives among the actual negatives. The false positive rate of the classifier is calculated as,

$$\hat{\gamma} = \frac{\#FP_k}{\#TN_k + \#FP_k}$$

Where:

- $\#TN_k$ is the number of true negatives, and
- $\#FP_k$ is the number of false positives.

Creating <factor>

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

See Also

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zerooneloss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zerooneloss.factor()`

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data    = iris,
  family  = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
predicted <- factor(
  as.numeric(
    predict(model, type = "response") > 0.5
  ),
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 4) evaluate class-wise performance
# using False Positive Rate

# 4.1) unweighted False Positive Rate
fpr(
  actual    = actual,
  predicted = predicted
)

# 4.2) weighted False Positive Rate
weighted.fpr(
  actual    = actual,
  predicted = predicted,
  w         = iris$Petal.Length/mean(iris$Petal.Length)
)
```

```

# 5) evaluate overall performance
# using micro-averaged False Positive Rate
cat(
  "Micro-averaged False Positive Rate", fpr(
    actual = actual,
    predicted = predicted,
    micro = TRUE
  ),
  "Micro-averaged False Positive Rate (weighted)", weighted.fpr(
    actual = actual,
    predicted = predicted,
    w = iris$Petal.Length/mean(iris$Petal.Length),
    micro = TRUE
  ),
  sep = "\n"
)

```

huberloss.numeric *Huber Loss*

Description

The `huberloss()`-function computes the simple and weighted **huber loss** between the predicted and observed `<numeric>` vectors. The `weighted.huberloss()` function computes the weighted Huber Loss.

Usage

```

## S3 method for class 'numeric'
huberloss(actual, predicted, delta = 1, ...)

## S3 method for class 'numeric'
weighted.huberloss(actual, predicted, w, delta = 1, ...)

## Generic S3 method
huberloss(
  actual,
  predicted,
  delta = 1,
  ...
)

## Generic S3 method
weighted.huberloss(
  actual,
  predicted,
  w,
  delta = 1,
  ...
)

```

Arguments

| | |
|-----------|--|
| actual | A <numeric>-vector of length n . The observed (continuous) response variable. |
| predicted | A <numeric>-vector of length n . The estimated (continuous) response variable. |
| delta | A <numeric>-vector of length 1 (default: 1). The threshold value for switch between functions (see calculation). |
| ... | Arguments passed into other methods. |
| w | A <numeric>-vector of length n . The weight assigned to each observation in the data. |

Value

A <numeric> vector of length 1.

Definition

The metric is calculated as follows,

$$\frac{1}{2}(y - v)^2 \text{ for } |y - v| \leq \delta$$

and

$$\delta|y - v| - \frac{1}{2}\delta^2 \text{ for otherwise}$$

where y and v are the actual and predicted values respectively. If w is not **NULL**, then all values are aggregated using the weights.

See Also

Other Regression: `ccc.numeric()`, `mae.numeric()`, `mape.numeric()`, `mpe.numeric()`, `mse.numeric()`, `pinball.numeric()`, `rae.numeric()`, `rmse.numeric()`, `rmsle.numeric()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroone.loss.factor()`

Examples

```
# 1) fit a linear
# regression
model <- lm(
  mpg ~ .,
  data = mtcars
```

```

)

# 1.1) define actual
# and predicted values
# to measure performance
actual <- mtcars$mpg
predicted <- fitted(model)

# 2) calculate the metric
# with delta 0.5
huberloss(
  actual = actual,
  predicted = predicted,
  delta = 0.5
)

# 3) calculate weighted
# metric using arbitrary weights
w <- rbeta(
  n = 1e3,
  shape1 = 10,
  shape2 = 2
)

huberloss(
  actual = actual,
  predicted = predicted,
  delta = 0.5,
  w = w
)

```

jaccard.factor

Jaccard Index

Description

The `jaccard()`-function computes the **Jaccard Index**, also known as the Intersection over Union, between two vectors of predicted and observed `factor()` values. The `weighted.jaccard()` function computes the weighted Jaccard Index.

Usage

```

## S3 method for class 'factor'
jaccard(actual, predicted, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
weighted.jaccard(actual, predicted, w, micro = NULL, na.rm = TRUE, ...)

```



```
## S3 method for class 'cmatrix'
jaccard(x, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
csi(actual, predicted, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
weighted.csi(actual, predicted, w, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'cmatrix'
csi(x, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
tscore(actual, predicted, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
weighted.tscore(actual, predicted, w, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'cmatrix'
tscore(x, micro = NULL, na.rm = TRUE, ...)

## Generic S3 method
jaccard(
  ...,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
csi(
  ...,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
tscore(
  ...,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
weighted.jaccard(
  ...,
  w,
  micro = NULL,
  na.rm = TRUE
)
```

```

)

## Generic S3 method
weighted.csi(
  ...,
  w,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
weighted.tscore(
  ...,
  w,
  micro = NULL,
  na.rm = TRUE
)

```

Arguments

| | |
|-----------|--|
| actual | A vector of <factor> values of length n , and k levels. |
| predicted | A vector of <factor> values of length n , and k levels. |
| micro | A <logical> -value of length 1 (default: NULL). If TRUE it returns the micro average across all k classes, if FALSE it returns the macro average. |
| na.rm | A <logical> value of length 1 (default: TRUE). If TRUE , NA values are removed from the computation. This argument is only relevant when <code>micro != NULL</code> . When <code>na.rm = TRUE</code> , the computation corresponds to <code>sum(c(1, 2, NA), na.rm = TRUE) / length(na.omit(c(1, 2, NA)))</code> . When <code>na.rm = FALSE</code> , the computation corresponds to <code>sum(c(1, 2, NA), na.rm = TRUE) / length(c(1, 2, NA))</code> . |
| ... | Arguments passed into other methods |
| w | A <numeric> -vector of length n . NULL by default. |
| x | A confusion matrix created <code>cmatrix()</code> . |

Value

If `micro` is [NULL](#) (the default), a named [<numeric>](#)-vector of [length](#) k
 If `micro` is [TRUE](#) or [FALSE](#), a [<numeric>](#)-vector of [length](#) 1

Definition

The metric is calculated for each class k as follows,

$$\frac{\#TP_k}{\#TP_k + \#FP_k + \#FN_k}$$

Where $\#TP_k$, $\#FP_k$, and $\#FN_k$ represent the number of true positives, false positives, and false negatives for each class k , respectively.

Creating <factor>

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

See Also

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zeroone.loss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroone.loss.factor()`

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data    = iris,
  family  = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
predicted <- factor(
  as.numeric(
    predict(model, type = "response") > 0.5
  ),
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 4) evaluate class-wise performance
# using Jaccard Index

# 4.1) unweighted Jaccard Index
jaccard(
  actual    = actual,
  predicted = predicted
)

# 4.2) weighted Jaccard Index
weighted.jaccard(
  actual    = actual,
  predicted = predicted,
  w         = iris$Petal.Length/mean(iris$Petal.Length)
)
```

```

# 5) evaluate overall performance
# using micro-averaged Jaccard Index
cat(
  "Micro-averaged Jaccard Index", jaccard(
    actual = actual,
    predicted = predicted,
    micro = TRUE
  ),
  "Micro-averaged Jaccard Index (weighted)", weighted.jaccard(
    actual = actual,
    predicted = predicted,
    w = iris$Petal.Length/mean(iris$Petal.Length),
    micro = TRUE
  ),
  sep = "\n"
)

```

| | |
|----------------|-----------------|
| logloss.factor | <i>Log Loss</i> |
|----------------|-----------------|

Description

The `logloss()` function computes the **Log Loss** between observed classes (as a `<factor>`) and their predicted probability distributions (a `<numeric>` matrix). The `weighted.logloss()` function is the weighted version, applying observation-specific weights.

Usage

```

## S3 method for class 'factor'
logloss(actual, response, normalize = TRUE, ...)

## S3 method for class 'factor'
weighted.logloss(actual, response, w, normalize = TRUE, ...)

## S3 method for class 'integer'
logloss(actual, response, normalize = TRUE, ...)

## S3 method for class 'integer'
weighted.logloss(actual, response, w, normalize = TRUE, ...)

## Generic S3 method
logloss(
  actual,
  response,
  normalize = TRUE,
  ...
)

```

```
## Generic S3 method
weighted.logloss(
  actual,
  response,
  w,
  normalize = TRUE,
  ...
)
```

Arguments

| | |
|-----------|---|
| actual | A vector of <code><factor></code> with length n , and k levels |
| response | A $n \times k$ <code><numeric></code> -matrix of predicted probabilities. The i -th row should sum to 1 (i.e., a valid probability distribution over the k classes). The first column corresponds to the first factor level in <code>actual</code> , the second column to the second factor level, and so on. |
| normalize | A <code><logical></code> -value (default: <code>TRUE</code>). If <code>TRUE</code> , the mean cross-entropy across all observations is returned; otherwise, the sum of cross-entropies is returned. |
| ... | Arguments passed into other methods |
| w | A <code><numeric></code> -vector of length n . <code>NULL</code> by default |

Value

A `<numeric>`-vector of length 1

Definition

$$H(p, response) = - \sum_i \sum_j y_{ij} \log_2(response_{ij})$$

where:

- y_{ij} is the actual-values, where $y_{ij} = 1$ if the i -th sample belongs to class j , and 0 otherwise.
- $response_{ij}$ is the estimated probability for the i -th sample belonging to class j .

Creating `<factor>`

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
```

```
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

See Also

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zeroone.loss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroone.loss.factor()`

Examples

```
# 1) Recode the iris data set to a binary classification problem
#   Here, the positive class ("Virginica") is coded as 1,
#   and the rest ("Others") is coded as 0.
iris$species_num <- as.numeric(iris$Species == "virginica")
```

```
# 2) Fit a logistic regression model predicting species_num from Sepal.Length & Sepal.Width
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data = iris,
  family = binomial(link = "logit")
)
```

```

)

# 3) Generate predicted classes: "Virginica" vs. "Others"
predicted <- factor(
  as.numeric(predict(model, type = "response") > 0.5),
  levels = c(1, 0),
  labels = c("Virginica", "Others")
)

# 3.1) Generate actual classes
actual <- factor(
  x      = iris$species_num,
  levels = c(1, 0),
  labels = c("Virginica", "Others")
)

# For Log Loss, we need predicted probabilities for each class.
# Since it's a binary model, we create a 2-column matrix:
# 1st column = P("Virginica")
# 2nd column = P("Others") = 1 - P("Virginica")
predicted_probs <- predict(model, type = "response")
response_matrix <- cbind(predicted_probs, 1 - predicted_probs)

# 4) Evaluate unweighted Log Loss
# 'logloss' takes (actual, response_matrix, normalize=TRUE/FALSE).
# The factor 'actual' must have the positive class (Virginica) as its first level.
unweighted_LogLoss <- logloss(
  actual      = actual,          # factor
  response    = response_matrix, # numeric matrix of probabilities
  normalize   = TRUE            # normalize = TRUE
)

# 5) Evaluate weighted Log Loss
# We introduce a weight vector, for example:
weights <- iris$Petal.Length / mean(iris$Petal.Length)
weighted_LogLoss <- weighted.logloss(
  actual      = actual,
  response    = response_matrix,
  w           = weights,
  normalize   = TRUE
)

# 6) Print Results
cat(
  "Unweighted Log Loss:", unweighted_LogLoss,
  "Weighted Log Loss:", weighted_LogLoss,
  sep = "\n"
)

```


Description

The `mae()`-function computes the **mean absolute error** between the observed and predicted `<numeric>` vectors. The `weighted.mae()` function computes the weighted mean absolute error.

Usage

```
## S3 method for class 'numeric'
mae(actual, predicted, ...)

## S3 method for class 'numeric'
weighted.mae(actual, predicted, w, ...)

## Generic S3 method
mae(
  actual,
  predicted,
  ...
)

## Generic S3 method
weighted.mae(
  actual,
  predicted,
  w,
  ...
)
```

Arguments

| | |
|------------------------|--|
| <code>actual</code> | A <code><numeric></code> -vector of <code>length n</code> . The observed (continuous) response variable. |
| <code>predicted</code> | A <code><numeric></code> -vector of <code>length n</code> . The estimated (continuous) response variable. |
| <code>...</code> | Arguments passed into other methods. |
| <code>w</code> | A <code><numeric></code> -vector of <code>length n</code> . The weight assigned to each observation in the data. |

Value

A `<numeric>` vector of `length 1`.

Definition

The metric is calculated as follows,

$$\frac{\sum_i^n |y_i - v_i|}{n}$$

See Also

Other Regression: `ccc.numeric()`, `huberloss.numeric()`, `mape.numeric()`, `mpe.numeric()`, `mse.numeric()`, `pinball.numeric()`, `rae.numeric()`, `rmse.numeric()`, `rmsle.numeric()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroone.loss.factor()`

Examples

```
# 1) fit a linear
# regression
model <- lm(
  mpg ~ .,
  data = mtcars
)

# 1.1) define actual
# and predicted values
# to measure performance
actual <- mtcars$mpg
predicted <- fitted(model)

# 2) evaluate in-sample model
# performance using Mean Absolute Error (MAE)
cat(
  "Mean Absolute Error", mae(
    actual = actual,
    predicted = predicted,
  ),
  "Mean Absolute Error (weighted)", weighted.mae(
    actual = actual,
    predicted = predicted,
    w = mtcars$mpg/mean(mtcars$mpg)
  ),
  sep = "\n"
)
```

Description

The `mape()`-function computes the **mean absolute percentage error** between the observed and predicted `<numeric>` vectors. The `weighted.mape()` function computes the weighted mean absolute percentage error.

Usage

```
## S3 method for class 'numeric'
mape(actual, predicted, ...)

## S3 method for class 'numeric'
weighted.mape(actual, predicted, w, ...)

## Generic S3 method
mape(
  actual,
  predicted,
  ...
)

## Generic S3 method
weighted.mape(
  actual,
  predicted,
  w,
  ...
)
```

Arguments

| | |
|------------------------|--|
| <code>actual</code> | A <code><numeric></code> -vector of length n . The observed (continuous) response variable. |
| <code>predicted</code> | A <code><numeric></code> -vector of length n . The estimated (continuous) response variable. |
| <code>...</code> | Arguments passed into other methods. |
| <code>w</code> | A <code><numeric></code> -vector of length n . The weight assigned to each observation in the data. |

Value

A `<numeric>` vector of **length** 1.

Definition

The metric is calculated as,

$$\frac{1}{n} \sum_i^n \frac{|y_i - v_i|}{|y_i|}$$

See Also

Other Regression: `ccc.numeric()`, `huberloss.numeric()`, `mae.numeric()`, `mpe.numeric()`, `mse.numeric()`, `pinball.numeric()`, `rae.numeric()`, `rmse.numeric()`, `rmsle.numeric()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroone.loss.factor()`

Examples

```
# 1) fit a linear
# regression
model <- lm(
  mpg ~ .,
  data = mtcars
)

# 1.1) define actual
# and predicted values
# to measure performance
actual <- mtcars$mpg
predicted <- fitted(model)

# 2) evaluate in-sample model
# performance using Mean Absolute Percentage Error (MAPE)
cat(
  "Mean Absolute Percentage Error", mape(
    actual = actual,
    predicted = predicted,
  ),
  "Mean Absolute Percentage Error (weighted)", weighted.mape(
    actual = actual,
    predicted = predicted,
    w = mtcars$mpg/mean(mtcars$mpg)
  ),
  sep = "\n"
)
```

Description

The `mcc()`-function computes the **Matthews Correlation Coefficient** (MCC), also known as the ϕ -coefficient, between two vectors of predicted and observed `factor()` values. The `weighted.mcc()` function computes the weighted Matthews Correlation Coefficient.

Usage

```
## S3 method for class 'factor'
mcc(actual, predicted, ...)

## S3 method for class 'factor'
weighted.mcc(actual, predicted, w, ...)

## S3 method for class 'cmatrix'
mcc(x, ...)

## S3 method for class 'factor'
phi(actual, predicted, ...)

## S3 method for class 'factor'
weighted.phi(actual, predicted, w, ...)

## S3 method for class 'cmatrix'
phi(x, ...)

## Generic S3 method
mcc(...)

## Generic S3 method
weighted.mcc(
  ...,
  w
)

## Generic S3 method
phi(...)

## Generic S3 method
weighted.phi(
  ...,
  w
)
```

Arguments

| | |
|------------------------|---|
| <code>actual</code> | A vector of <code><factor></code> with length n , and k levels |
| <code>predicted</code> | A vector of <code><factor></code> with length n , and k levels |
| <code>...</code> | Arguments passed into other methods |

w A <numeric>-vector of length n . NULL by default
 x A confusion matrix created `cmatrix()`

Value

A <numeric>-vector of length 1

Definition

The metric is calculated as follows,

$$\frac{\#TP \times \#TN - \#FP \times \#FN}{\sqrt{(\#TP + \#FP)(\#TP + \#FN)(\#TN + \#FP)(\#TN + \#FN)}}$$

Creating <factor>

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

See Also

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zerooneloss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zerooneloss.factor()`

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data = iris,
  family = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
predicted <- factor(
  as.numeric(
    predict(model, type = "response") > 0.5
  ),
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)
```

```
# 4) evaluate performance
# using Matthews Correlation Coefficient
cat(
  "Matthews Correlation Coefficient", mcc(
    actual = actual,
    predicted = predicted
  ),
  "Matthews Correlation Coefficient (weighted)", weighted.mcc(
    actual = actual,
    predicted = predicted,
    w = iris$Petal.Length/mean(iris$Petal.Length)
  ),
  sep = "\n"
)
```

mpe.numeric

Mean Percentage Error

Description

The `mpe()`-function computes the **mean percentage error** between the observed and predicted `<numeric>` vectors. The `weighted.mpe()` function computes the weighted mean percentage error.

Usage

```
## S3 method for class 'numeric'
mpe(actual, predicted, ...)

## S3 method for class 'numeric'
weighted.mpe(actual, predicted, w, ...)

## Generic S3 method
mpe(
  actual,
  predicted,
  ...
)

## Generic S3 method
weighted.mpe(
  actual,
  predicted,
  w,
  ...
)
```


Arguments

| | |
|-----------|---|
| actual | A <numeric>-vector of length n . The observed (continuous) response variable. |
| predicted | A <numeric>-vector of length n . The estimated (continuous) response variable. |
| ... | Arguments passed into other methods. |
| w | A <numeric>-vector of length n . The weight assigned to each observation in the data. |

Value

A <numeric> vector of length 1.

Definition

The metric is calculated as,

$$\frac{1}{n} \sum_i^n \frac{y_i - v_i}{y_i}$$

Where y_i and v_i are the actual and predicted values respectively.

See Also

Other Regression: [ccc.numeric\(\)](#), [huberloss.numeric\(\)](#), [mae.numeric\(\)](#), [mape.numeric\(\)](#), [mse.numeric\(\)](#), [pinball.numeric\(\)](#), [rae.numeric\(\)](#), [rmse.numeric\(\)](#), [rmsle.numeric\(\)](#), [rrmse.numeric\(\)](#), [rrse.numeric\(\)](#), [rsq.numeric\(\)](#), [smape.numeric\(\)](#)

Other Supervised Learning: [ROC.factor\(\)](#), [accuracy.factor\(\)](#), [baccuracy.factor\(\)](#), [ccc.numeric\(\)](#), [ckappa.factor\(\)](#), [cmatrix.factor\(\)](#), [dor.factor\(\)](#), [entropy.matrix\(\)](#), [fbeta.factor\(\)](#), [fdr.factor\(\)](#), [fer.factor\(\)](#), [fpr.factor\(\)](#), [huberloss.numeric\(\)](#), [jaccard.factor\(\)](#), [logloss.factor\(\)](#), [mae.numeric\(\)](#), [mape.numeric\(\)](#), [mcc.factor\(\)](#), [mse.numeric\(\)](#), [nlr.factor\(\)](#), [npv.factor\(\)](#), [pinball.numeric\(\)](#), [plr.factor\(\)](#), [pr.auc.matrix\(\)](#), [prROC.factor\(\)](#), [precision.factor\(\)](#), [rae.numeric\(\)](#), [recall.factor\(\)](#), [rmse.numeric\(\)](#), [rmsle.numeric\(\)](#), [roc.auc.matrix\(\)](#), [rrmse.numeric\(\)](#), [rrse.numeric\(\)](#), [rsq.numeric\(\)](#), [smape.numeric\(\)](#), [specificity.factor\(\)](#), [zerooneloss.factor\(\)](#)

Examples

```
# 1) fit a linear
# regression
model <- lm(
  mpg ~ .,
  data = mtcars
)

# 1.1) define actual
# and predicted values
# to measure performance
actual <- mtcars$mpg
predicted <- fitted(model)
```

```

# 2) evaluate in-sample model
# performance using Mean Percentage Error (MPE)
cat(
  "Mean Percentage Error", mpe(
    actual = actual,
    predicted = predicted,
  ),
  "Mean Percentage Error (weighted)", weighted.mpe(
    actual = actual,
    predicted = predicted,
    w = mtcars$mpg/mean(mtcars$mpg)
  ),
  sep = "\n"
)

```

mse.numeric

Mean Squared Error

Description

The `mse()`-function computes the **mean squared error** between the observed and predicted `<numeric>` vectors. The `weighted.mse()` function computes the weighted mean squared error.

Usage

```

## S3 method for class 'numeric'
mse(actual, predicted, ...)

## S3 method for class 'numeric'
weighted.mse(actual, predicted, w, ...)

## Generic S3 method
mse(
  actual,
  predicted,
  ...
)

## Generic S3 method
weighted.mse(
  actual,
  predicted,
  w,
  ...
)

```

Arguments

| | |
|-----------|---|
| actual | A <numeric>-vector of length n . The observed (continuous) response variable. |
| predicted | A <numeric>-vector of length n . The estimated (continuous) response variable. |
| ... | Arguments passed into other methods. |
| w | A <numeric>-vector of length n . The weight assigned to each observation in the data. |

Value

A <numeric> vector of length 1.

Definition

The metric is calculated as,

$$\frac{1}{n} \sum_i^n (y_i - v_i)^2$$

Where y_i and v_i are the actual and predicted values respectively.

See Also

Other Regression: `ccc.numeric()`, `huberloss.numeric()`, `mae.numeric()`, `mape.numeric()`, `mpe.numeric()`, `pinball.numeric()`, `rae.numeric()`, `rmse.numeric()`, `rmsle.numeric()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zerooneloss.factor()`

Examples

```
# 1) fit a linear
# regression
model <- lm(
  mpg ~ .,
  data = mtcars
)

# 1.1) define actual
# and predicted values
# to measure performance
actual <- mtcars$mpg
predicted <- fitted(model)
```

```

# 2) evaluate in-sample model
# performance using Mean Squared Error (MSE)
cat(
  "Mean Squared Error", mse(
    actual = actual,
    predicted = predicted,
  ),
  "Mean Squared Error (weighted)", weighted.mse(
    actual = actual,
    predicted = predicted,
    w = mtcars$mpg/mean(mtcars$mpg)
  ),
  sep = "\n"
)

```

nlr.factor

Negative Likelihood Ratio

Description

A generic function for the **negative likelihood ratio** in classification tasks. Use `weighted.nlr()` weighted **negative likelihood ratio**.

Usage

```

## S3 method for class 'factor'
nlr(actual, predicted, ...)

## S3 method for class 'factor'
weighted.nlr(actual, predicted, w, ...)

## S3 method for class 'cmatrix'
nlr(x, ...)

## Generic S3 method
nlr(...)

## Generic S3 method
weighted.nlr(
  ...,
  w
)

```

Arguments

`actual` A vector of `<factor>` values of `length n`, and `k` levels.
`predicted` A vector of `<factor>` values of `length n`, and `k` levels.

... Arguments passed into other methods
 w A <numeric>-vector of length n . NULL by default.
 x A confusion matrix created `cmatrix()`.

Value

If `micro` is NULL (the default), a named <numeric>-vector of length k

If `micro` is TRUE or FALSE, a <numeric>-vector of length 1

Definition

Let $\hat{\alpha} \in [0, \infty]$ be the likelihood of a negative outcome. The **negative likelihood ratio** of the classifier is calculated as,

$$\hat{\alpha} = \frac{1 - \frac{\#TP}{\#TP + \#FN}}{\frac{\#TN}{\#TN + \#FP}}$$

Where:

- $\frac{\#TP}{\#TP + \#FN}$ is the sensitivity, or true positive rate
- $\frac{\#TN}{\#TN + \#FP}$ is the specificity, or true negative rate

Creating <factor>

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
```

```

x = sample(x = c(1, 3), size = 10, replace = TRUE),
levels = c(1, 2, 3),
labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C

```

In both cases, $k = 3$, determined indirectly by the levels argument.

See Also

The `plr()`-function for the Positive Likelihood Ratio (LR+)

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zeroone.loss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroone.loss.factor()`

Examples

```

# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data = iris,
  family = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
predicted <- factor(
  as.numeric(
    predict(model, type = "response") > 0.5
  )
)

```

```

    ),
    levels = c(1,0),
    labels = c("Virginica", "Others")
  )

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 4) evaluate model performance
# with class-wise negative likelihood ratios
cat("Negative Likelihood Ratio", sep = "\n")
nlr(
  actual = actual,
  predicted = predicted
)

cat("Negative Likelihood Ratio (weighted)", sep = "\n")
weighted.nlr(
  actual = actual,
  predicted = predicted,
  w = iris$Petal.Length/mean(iris$Petal.Length)
)

```

npv.factor

Negative Predictive Value

Description

The `npv()`-function computes the **negative predictive value**, also known as the True Negative Predictive Value, between two vectors of predicted and observed `factor()` values. The `weighted.npv()` function computes the weighted negative predictive value.

Usage

```

## S3 method for class 'factor'
npv(actual, predicted, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
weighted.npv(actual, predicted, w, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'cmatrix'
npv(x, micro = NULL, na.rm = TRUE, ...)

npv(...)

```

```
weighted.npv(...)
```

Arguments

| | |
|-----------|---|
| actual | A vector of <factor> values of length n , and k levels. |
| predicted | A vector of <factor> values of length n , and k levels. |
| micro | A <logical> -value of length 1 (default: NULL). If TRUE it returns the micro average across all k classes, if FALSE it returns the macro average. |
| na.rm | A <logical> value of length 1 (default: TRUE). If TRUE , NA values are removed from the computation. This argument is only relevant when micro != NULL . When na.rm = TRUE , the computation corresponds to <code>sum(c(1, 2, NA), na.rm = TRUE) / length(na.omit(c(1, 2, NA)))</code> . When na.rm = FALSE , the computation corresponds to <code>sum(c(1, 2, NA), na.rm = TRUE) / length(c(1, 2, NA))</code> . |
| ... | Arguments passed into other methods |
| w | A <numeric> -vector of length n . NULL by default. |
| x | A confusion matrix created <code>cmatrix()</code> . |

Value

If **micro** is **NULL** (the default), a named **<numeric>**-vector of **length** k

If **micro** is **TRUE** or **FALSE**, a **<numeric>**-vector of **length** 1

Definition

The metric is calculated for each class k as follows,

$$\frac{\#TN_k}{\#TN_k + \#FN_k}$$

Where $\#TN_k$ and $\#FN_k$ are the number of true negatives and false negatives, respectively, for each class k .

Creating **<factor>**

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
```



```
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

See Also

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zeroone.loss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroone.loss.factor()`

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
```

```
    formula = species_num ~ Sepal.Length + Sepal.Width,
    data     = iris,
    family   = binomial(
      link = "logit"
    )
  )
)

# 3) generate predicted
# classes
predicted <- factor(
  as.numeric(
    predict(model, type = "response") > 0.5
  ),
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 4) evaluate class-wise performance
# using Negative Predictive Value

# 4.1) unweighted Negative Predictive Value
npv(
  actual     = actual,
  predicted  = predicted
)

# 4.2) weighted Negative Predictive Value
weighted.npv(
  actual     = actual,
  predicted  = predicted,
  w          = iris$Petal.Length/mean(iris$Petal.Length)
)

# 5) evaluate overall performance
# using micro-averaged Negative Predictive Value
cat(
  "Micro-averaged Negative Predictive Value", npv(
    actual     = actual,
    predicted  = predicted,
    micro      = TRUE
  ),
  "Micro-averaged Negative Predictive Value (weighted)", weighted.npv(
    actual     = actual,
    predicted  = predicted,
    w          = iris$Petal.Length/mean(iris$Petal.Length),
```

```

    micro      = TRUE
  ),
  sep = "\n"
)

```

obesity

Obesity Levels Dataset

Description

This dataset is used to estimate obesity levels based on eating habits and physical condition. The data originates from the UCI Machine Learning Repository and has been preprocessed to include both predictors and a target variable.

Usage

```
data(obesity)
```

Format

A list with two components:

features A data frame containing various predictors related to eating habits, physical condition, and lifestyle.

target A list with two elements: regression (weight in kilograms) and class (obesity level classification).

Details

The dataset is provided as a list with two components:

features A data frame containing various predictors related to lifestyle, eating habits, and physical condition. The variables include:

age The age of the individual in years.

height The height of the individual in meters.

family_history_with_overweight Binary variable indicating whether the individual has a family history of overweight (1 = yes, 0 = no).

fave Binary variable indicating whether the individual frequently consumes high-calorie foods (1 = yes, 0 = no).

fcvc The frequency of consumption of vegetables in meals.

ncp The number of main meals consumed per day.

caec Categorical variable indicating the frequency of consumption of food between meals. Typical levels include "no", "sometimes", "frequently", and "always".

smoke Binary variable indicating whether the individual smokes (1 = yes, 0 = no).

ch2o Daily water consumption (typically in liters).

scc Binary variable indicating whether the individual monitors calorie consumption (1 = yes, 0 = no).

faf The frequency of physical activity.

tue The time spent using electronic devices (e.g., screen time in hours).

calc Categorical variable indicating the frequency of alcohol consumption. Typical levels include "no", "sometimes", "frequently", and "always".

male Binary variable indicating the gender of the individual (1 = male, 0 = female).

target A list containing two elements:

regression A numeric vector representing the weight of the individual (used as the regression target).

class A factor indicating the obesity level classification. The levels are derived from the original nobeyesdad variable in the dataset.

Source

<https://archive.ics.uci.edu/dataset/544/estimation+of+obesity+levels+based+on+eating+habits+and+physical+condition>

openmp.on

Use OpenMP

Description

This function allows you to enable or disable the use of OpenMP for parallelizing computations.

Usage

```
## enable OpenMP
openmp.on()
```

```
## disable OpenMP
openmp.off()
```

```
## set number of threads
openmp.threads(threads)
```

Arguments

threads A positive [<integer>](#)-value (Default: None). If threads is missing, the `openmp.threads()` returns the number of available threads. If [NULL](#) all available threads will be used.

Value

If OpenMP is unavailable, the function returns [NULL](#).

If OpenMP is unavailable, the function returns [NULL](#).

If OpenMP is unavailable, the function returns [NULL](#).

Examples

```
## Not run:
## enable OpenMP
SLmetrics::openmp.on()

## disable OpenMP
SLmetrics::openmp.off()

## available threads
SLmetrics::openmp.threads()

## set number of threads
SLmetrics::openmp.threads(2)

## End(Not run)
```

| | |
|-----------------|---------------------|
| pinball.numeric | <i>Pinball Loss</i> |
|-----------------|---------------------|

Description

The `pinball()`-function computes the **pinball loss** between the observed and predicted `<numeric>` vectors. The `weighted.pinball()` function computes the weighted Pinball Loss.

Usage

```
## S3 method for class 'numeric'
pinball(actual, predicted, alpha = 0.5, deviance = FALSE, ...)

## S3 method for class 'numeric'
weighted.pinball(actual, predicted, w, alpha = 0.5, deviance = FALSE, ...)

## Generic S3 method
pinball(
  actual,
  predicted,
  alpha = 0.5,
  deviance = FALSE,
  ...
)

## Generic S3 method
weighted.pinball(
  actual,
  predicted,
  w,
```

```

alpha    = 0.5,
deviance = FALSE,
...
)

```

Arguments

| | |
|-----------|--|
| actual | A <numeric>-vector of length n . The observed (continuous) response variable. |
| predicted | A <numeric>-vector of length n . The estimated (continuous) response variable. |
| alpha | A <numeric>-value of length 1 (default: 0.5). The slope of the pinball loss function. |
| deviance | A <logical>-value of length 1 (default: FALSE). If TRUE the function returns the D^2 loss. |
| ... | Arguments passed into other methods. |
| w | A <numeric>-vector of length n . The weight assigned to each observation in the data. |

Value

A <numeric> vector of length 1.

Definition

The metric is calculated as,

$$\text{PinballLoss}_{\text{unweighted}} = \frac{1}{n} \sum_{i=1}^n [\alpha \cdot \max(0, y_i - \hat{y}_i) - (1 - \alpha) \cdot \max(0, \hat{y}_i - y_i)]$$

where y_i is the actual value, \hat{y}_i is the predicted value and α is the quantile level.

See Also

Other Regression: [ccc.numeric\(\)](#), [huberloss.numeric\(\)](#), [mae.numeric\(\)](#), [mape.numeric\(\)](#), [mpe.numeric\(\)](#), [mse.numeric\(\)](#), [rae.numeric\(\)](#), [rmse.numeric\(\)](#), [rmsle.numeric\(\)](#), [rrmse.numeric\(\)](#), [rrse.numeric\(\)](#), [rsq.numeric\(\)](#), [smape.numeric\(\)](#)

Other Supervised Learning: [ROC.factor\(\)](#), [accuracy.factor\(\)](#), [baccuracy.factor\(\)](#), [ccc.numeric\(\)](#), [ckappa.factor\(\)](#), [cmatrix.factor\(\)](#), [dor.factor\(\)](#), [entropy.matrix\(\)](#), [fbeta.factor\(\)](#), [fdr.factor\(\)](#), [fer.factor\(\)](#), [fpr.factor\(\)](#), [huberloss.numeric\(\)](#), [jaccard.factor\(\)](#), [logloss.factor\(\)](#), [mae.numeric\(\)](#), [mape.numeric\(\)](#), [mcc.factor\(\)](#), [mpe.numeric\(\)](#), [mse.numeric\(\)](#), [nlr.factor\(\)](#), [npv.factor\(\)](#), [plr.factor\(\)](#), [pr.auc.matrix\(\)](#), [prROC.factor\(\)](#), [precision.factor\(\)](#), [rae.numeric\(\)](#), [recall.factor\(\)](#), [rmse.numeric\(\)](#), [rmsle.numeric\(\)](#), [roc.auc.matrix\(\)](#), [rrmse.numeric\(\)](#), [rrse.numeric\(\)](#), [rsq.numeric\(\)](#), [smape.numeric\(\)](#), [specificity.factor\(\)](#), [zeroonloss.factor\(\)](#)

Examples

```

# 1) fit a linear
# regression
model <- lm(
  mpg ~ .,
  data = mtcars
)

# 1.1) define actual
# and predicted values
# to measure performance
actual <- mtcars$mpg
predicted <- fitted(model)

# 2) evaluate in-sample model
# performance using Pinball Loss
cat(
  "Pinball Loss", pinball(
    actual = actual,
    predicted = predicted,
  ),
  "Pinball Loss (weighted)", weighted.pinball(
    actual = actual,
    predicted = predicted,
    w = mtcars$mpg/mean(mtcars$mpg)
  ),
  sep = "\n"
)

```

 plr.factor

Positive Likelihood Ratio

Description

A generic function for the **positive likelihood ratio** in classification tasks. Use `weighted.plr()` weighted **positive likelihood ratio**.

Usage

```

## S3 method for class 'factor'
plr(actual, predicted, ...)

## S3 method for class 'factor'
weighted.plr(actual, predicted, w, ...)

## S3 method for class 'cmatrix'
plr(x, ...)

## Generic S3 method

```

```
plr(...)

## Generic S3 method
weighted.plr(
  ...,
  w
)
```

Arguments

| | |
|-----------|---|
| actual | A vector of <factor> values of length n , and k levels. |
| predicted | A vector of <factor> values of length n , and k levels. |
| ... | Arguments passed into other methods |
| w | A <numeric> -vector of length n . NULL by default. |
| x | A confusion matrix created <code>cmatrix()</code> . |

Value

If `micro` is **NULL** (the default), a named **<numeric>**-vector of **length** k

If `micro` is **TRUE** or **FALSE**, a **<numeric>**-vector of **length** 1

Definition

Let $\hat{\alpha} \in [0, \infty]$ be the likelihood of a positive outcome. The **positive likelihood ratio** of the classifier is calculated as,

$$\hat{\alpha} = \frac{\frac{\#TP}{\#TP+\#FN}}{1 - \frac{\#TN}{\#TN+\#FP}}$$

Where:

- $\frac{\#TP}{\#TP+\#FN}$ is the sensitivity, or true positive rate
- $\frac{\#TN}{\#TN+\#FP}$ is the specificity, or true negative rate

Creating **<factor>**

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
```



```
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

See Also

The `nlr()`-function for the Negative Likelihood Ratio (LR-)

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zeroone.loss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroone.loss.factor()`

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
```

```

model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data    = iris,
  family  = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
predicted <- factor(
  as.numeric(
    predict(model, type = "response") > 0.5
  ),
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 4) evaluate model performance
# with class-wise positive likelihood ratios
cat("Positive Likelihood Ratio", sep = "\n")
plr(
  actual    = actual,
  predicted = predicted
)

cat("Positive Likelihood Ratio (weighted)", sep = "\n")
weighted.plr(
  actual    = actual,
  predicted = predicted,
  w         = iris$Petal.Length/mean(iris$Petal.Length)
)

```

pr.auc.matrix

Area under the Precision-Recall Curve

Description

A generic function for the area under the Precision-Recall Curve. Use `weighted.pr.auc()` for the weighted area under the Precision-Recall Curve.

Usage

```
## S3 method for class 'matrix'
pr.auc(actual, response, micro = NULL, method = 0L, ...)

## S3 method for class 'matrix'
weighted.pr.auc(actual, response, w, micro = NULL, method = 0L, ...)

## Generic S3 method
pr.auc(
  actual,
  response,
  micro = NULL,
  method = 0,
  ...
)

## Generic S3 method
weighted.pr.auc(
  actual,
  response,
  w,
  micro = NULL,
  method = 0,
  ...
)
```

Arguments

| | |
|----------|--|
| actual | A vector of <factor> values of length n , and k levels. |
| response | A $n \times k$ <numeric>-matrix . The estimated response probabilities for each class k . |
| micro | A <logical> -value of length 1 (default: NULL). If TRUE it returns the micro average across all k classes, if FALSE it returns the macro average. |
| method | A <numeric> value (default: 0). Defines the underlying method of calculating the area under the curve. If 0 it is calculated using the trapezoid-method, if 1 it is calculated using the step-method. |
| ... | Arguments passed into other methods. |
| w | A <numeric> -vector of length n . NULL by default. |

Value

A [<numeric>](#) vector of [length](#) 1

Definition**Trapezoidal rule**

The **trapezoidal rule** approximates the integral of a function $f(x)$ between $x = a$ and $x = b$ using trapezoids formed between consecutive points. If we have points x_0, x_1, \dots, x_n (with $a = x_0 < x_1 < \dots < x_n = b$) and corresponding function values $f(x_0), f(x_1), \dots, f(x_n)$, the area under the curve A_T is approximated by:

$$A_T \approx \sum_{k=1}^n \frac{f(x_{k-1}) + f(x_k)}{2} [x_k - x_{k-1}].$$

Step-function method

The **step-function (rectangular) method** uses the value of the function at one endpoint of each subinterval to form rectangles. With the same partition x_0, x_1, \dots, x_n , the rectangular approximation A_S can be written as:

$$A_S \approx \sum_{k=1}^n f(x_{k-1}) [x_k - x_{k-1}].$$

See Also

Other Classification: [ROC.factor\(\)](#), [accuracy.factor\(\)](#), [baccuracy.factor\(\)](#), [ckappa.factor\(\)](#), [cmatrix.factor\(\)](#), [dor.factor\(\)](#), [entropy.matrix\(\)](#), [fbeta.factor\(\)](#), [fdr.factor\(\)](#), [fer.factor\(\)](#), [fmi.factor\(\)](#), [fpr.factor\(\)](#), [jaccard.factor\(\)](#), [logloss.factor\(\)](#), [mcc.factor\(\)](#), [nlr.factor\(\)](#), [npv.factor\(\)](#), [plr.factor\(\)](#), [prROC.factor\(\)](#), [precision.factor\(\)](#), [recall.factor\(\)](#), [roc.auc.matrix\(\)](#), [specificity.factor\(\)](#), [zeroone.loss.factor\(\)](#)

Other Supervised Learning: [ROC.factor\(\)](#), [accuracy.factor\(\)](#), [baccuracy.factor\(\)](#), [ccc.numeric\(\)](#), [ckappa.factor\(\)](#), [cmatrix.factor\(\)](#), [dor.factor\(\)](#), [entropy.matrix\(\)](#), [fbeta.factor\(\)](#), [fdr.factor\(\)](#), [fer.factor\(\)](#), [fpr.factor\(\)](#), [huberloss.numeric\(\)](#), [jaccard.factor\(\)](#), [logloss.factor\(\)](#), [mae.numeric\(\)](#), [mape.numeric\(\)](#), [mcc.factor\(\)](#), [mpe.numeric\(\)](#), [mse.numeric\(\)](#), [nlr.factor\(\)](#), [npv.factor\(\)](#), [pinball.numeric\(\)](#), [plr.factor\(\)](#), [prROC.factor\(\)](#), [precision.factor\(\)](#), [rae.numeric\(\)](#), [recall.factor\(\)](#), [rmse.numeric\(\)](#), [rmsle.numeric\(\)](#), [roc.auc.matrix\(\)](#), [rrmse.numeric\(\)](#), [rrse.numeric\(\)](#), [rsq.numeric\(\)](#), [smape.numeric\(\)](#), [specificity.factor\(\)](#), [zeroone.loss.factor\(\)](#)

Other Classification: [ROC.factor\(\)](#), [accuracy.factor\(\)](#), [baccuracy.factor\(\)](#), [ckappa.factor\(\)](#), [cmatrix.factor\(\)](#), [dor.factor\(\)](#), [entropy.matrix\(\)](#), [fbeta.factor\(\)](#), [fdr.factor\(\)](#), [fer.factor\(\)](#), [fmi.factor\(\)](#), [fpr.factor\(\)](#), [jaccard.factor\(\)](#), [logloss.factor\(\)](#), [mcc.factor\(\)](#), [nlr.factor\(\)](#), [npv.factor\(\)](#), [plr.factor\(\)](#), [prROC.factor\(\)](#), [precision.factor\(\)](#), [recall.factor\(\)](#), [roc.auc.matrix\(\)](#), [specificity.factor\(\)](#), [zeroone.loss.factor\(\)](#)

Other Supervised Learning: [ROC.factor\(\)](#), [accuracy.factor\(\)](#), [baccuracy.factor\(\)](#), [ccc.numeric\(\)](#), [ckappa.factor\(\)](#), [cmatrix.factor\(\)](#), [dor.factor\(\)](#), [entropy.matrix\(\)](#), [fbeta.factor\(\)](#), [fdr.factor\(\)](#), [fer.factor\(\)](#), [fpr.factor\(\)](#), [huberloss.numeric\(\)](#), [jaccard.factor\(\)](#), [logloss.factor\(\)](#), [mae.numeric\(\)](#), [mape.numeric\(\)](#), [mcc.factor\(\)](#), [mpe.numeric\(\)](#), [mse.numeric\(\)](#), [nlr.factor\(\)](#), [npv.factor\(\)](#), [pinball.numeric\(\)](#), [plr.factor\(\)](#), [prROC.factor\(\)](#), [precision.factor\(\)](#), [rae.numeric\(\)](#), [recall.factor\(\)](#), [rmse.numeric\(\)](#), [rmsle.numeric\(\)](#), [roc.auc.matrix\(\)](#), [rrmse.numeric\(\)](#), [rrse.numeric\(\)](#), [rsq.numeric\(\)](#), [smape.numeric\(\)](#), [specificity.factor\(\)](#), [zeroone.loss.factor\(\)](#)

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data    = iris,
  family  = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
response <- predict(model, type = "response")

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 4) generate precision-recall
# data

# 4.1) calculate residual
# probability and store as matrix
response <- matrix(
  data = cbind(response, 1-response),
  nrow = length(actual)
)

# 4.2) calculate class-wise
# area under the curve
pr.auc(
  actual    = actual,
  response  = response
)

# 4.3) calculate class-wise
# weighted area under the curve
weighted.pr.auc(
  actual    = actual,
  response  = response,
```

```

    w      = iris$Petal.Length/mean(iris$Petal.Length)
  )

# 5) evaluate overall area under
# the curve
cat(
  "Micro-averaged area under the precision-recall curve", pr.auc(
    actual   = actual,
    response = response,
    micro    = TRUE
  ),
  "Micro-averaged area under the precision-recall curve (weighted)", weighted.pr.auc(
    actual   = actual,
    response = response,
    w        = iris$Petal.Length/mean(iris$Petal.Length),
    micro    = TRUE
  ),
  sep = "\n"
)

```

| | |
|------------------|------------------|
| precision.factor | <i>Precision</i> |
|------------------|------------------|

Description

A generic function for the **precision**. Use `weighted.fdr()` for the weighted **precision**.

Other names:

Positive Predictive Value

Usage

```

## S3 method for class 'factor'
precision(actual, predicted, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
weighted.precision(actual, predicted, w, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'cmatrix'
precision(x, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
ppv(actual, predicted, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
weighted.ppv(actual, predicted, w, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'cmatrix'
ppv(x, micro = NULL, na.rm = TRUE, ...)

```

```

## Generic S3 method
precision(
  ...,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
weighted.precision(
  ...,
  w,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
ppv(
  ...,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
weighted.ppv(
  ...,
  w,
  micro = NULL,
  na.rm = TRUE
)

```

Arguments

| | |
|-----------|--|
| actual | A vector of <code><factor></code> values of length n , and k levels. |
| predicted | A vector of <code><factor></code> values of length n , and k levels. |
| micro | A <code><logical></code> -value of length 1 (default: <code>NULL</code>). If <code>TRUE</code> it returns the micro average across all k classes, if <code>FALSE</code> it returns the macro average. |
| na.rm | A <code><logical></code> value of length 1 (default: <code>TRUE</code>). If <code>TRUE</code> , <code>NA</code> values are removed from the computation. This argument is only relevant when <code>micro != NULL</code> . When <code>na.rm = TRUE</code> , the computation corresponds to <code>sum(c(1, 2, NA), na.rm = TRUE) / length(na.omit(c(1, 2, NA)))</code> . When <code>na.rm = FALSE</code> , the computation corresponds to <code>sum(c(1, 2, NA), na.rm = TRUE) / length(c(1, 2, NA))</code> . |
| ... | Arguments passed into other methods |
| w | A <code><numeric></code> -vector of length n . <code>NULL</code> by default. |
| x | A confusion matrix created <code>cmatrix()</code> . |

Value

If `micro` is `NULL` (the default), a named `<numeric>`-vector of length `k`

If `micro` is `TRUE` or `FALSE`, a `<numeric>`-vector of length 1

Definition

Let $\hat{\pi} \in [0, 1]$ be the proportion of true positives among the predicted positives. The precision of the classifier is calculated as,

$$\hat{\pi} = \frac{\#TP_k}{\#TP_k + \#FP_k}$$

Where:

- $\#TP_k$ is the number of true positives, and
- $\#FP_k$ is the number of false positives.

Creating `<factor>`

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

See Also

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zerooneloss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zerooneloss.factor()`

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data = iris,
  family = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
predicted <- factor(
  as.numeric(
    predict(model, type = "response") > 0.5
  ),
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)
```

```

# 4) evaluate class-wise performance
# using Precision

# 4.1) unweighted Precision
precision(
  actual = actual,
  predicted = predicted
)

# 4.2) weighted Precision
weighted.precision(
  actual = actual,
  predicted = predicted,
  w = iris$Petal.Length/mean(iris$Petal.Length)
)

# 5) evaluate overall performance
# using micro-averaged Precision
cat(
  "Micro-averaged Precision", precision(
    actual = actual,
    predicted = predicted,
    micro = TRUE
  ),
  "Micro-averaged Precision (weighted)", weighted.precision(
    actual = actual,
    predicted = predicted,
    w = iris$Petal.Length/mean(iris$Petal.Length),
    micro = TRUE
  ),
  sep = "\n"
)

```

preorder

Preorder

Description

This function does a column-wise ordering permutation of [numeric](#) or [integer](#) matrix.

Usage

```

preorder(
  x,
  decreasing = FALSE,
  ...
)

```

Arguments

`x` a [numeric](#) or [integer](#) matrix to be sorted.
`decreasing` a [logical](#) value of [length](#) 1 (default: [FALSE](#)). If [TRUE](#) the matrix is returned in descending order.
... Arguments passed into other methods.

Value

A [matrix](#) with indices to the ordered values.

See Also

Other Tools: [auc.numeric\(\)](#), [cov.wt.matrix\(\)](#), [presort\(\)](#)

Examples

```
# 1) generate a 4x4 matrix
# with random values to be sorted
set.seed(1903)
X <- matrix(
  data = cbind(sample(16:1)),
  nrow = 4
)

# 2) sort matrix
# in decreasing order
presort(X)

# 3) get indices
# for sorted matrix
preorder(X)
```

presort

Presort

Description

This generic function does a column-wise sorting of a [numeric](#) or [integer](#) matrix.

Usage

```
presort(
  x,
  decreasing = FALSE,
  ...
)
```

Arguments

`x` a [numeric](#) or [integer](#) matrix to be sorted.

`decreasing` a [logical](#) value of [length](#) 1 (default: [FALSE](#)). If [TRUE](#) the matrix is returned in descending order.

`...` Arguments passed into other methods.

Value

A [matrix](#) with sorted rows.

See Also

Other Tools: [auc.numeric\(\)](#), [cov.wt.matrix\(\)](#), [preorder\(\)](#)

Examples

```
# 1) generate a 4x4 matrix
# with random values to be sorted
set.seed(1903)
X <- matrix(
  data = cbind(sample(16:1)),
  nrow = 4
)

# 2) sort matrix
# in decreasing order
presort(X)

# 3) get indices
# for sorted matrix
preorder(X)
```

```
prROC.factor
```

```
Precision-Recall Curve
```

Description

The [prROC\(\)](#)-function computes the [precision\(\)](#) and [recall\(\)](#) at thresholds provided by the *response*- or *thresholds*-vector. The function constructs a [data.frame\(\)](#) grouped by *k*-classes where each class is treated as a binary classification problem.

Usage

```
## S3 method for class 'factor'
prROC(actual, response, thresholds = NULL, presorted = FALSE, ...)

## S3 method for class 'factor'
weighted.prROC(actual, response, w, thresholds = NULL, presorted = FALSE, ...)
```

```
## Generic S3 method
prROC(
  actual,
  response,
  thresholds = NULL,
  presorted = FALSE,
  ...
)

## Generic S3 method
weighted.prROC(
  actual,
  response,
  w,
  thresholds = NULL,
  presorted = FALSE,
  ...
)
```

Arguments

| | |
|------------|---|
| actual | A vector of <factor> values of length n , and k levels. |
| response | A $n \times k$ <numeric>-matrix . The estimated response probabilities for each class k . |
| thresholds | An optional <numeric> vector of length n (default: NULL). |
| presorted | A <logical> -value length 1 (default: FALSE). If TRUE the input will not be sorted by threshold. |
| ... | Arguments passed into other methods. |
| w | A <numeric> -vector of length n . NULL by default. |

Value

A [data.frame](#) on the following form,

| | |
|-----------|---|
| threshold | <numeric> Thresholds used to determine recall() and precision() |
| level | <character> The level of the actual <factor> |
| label | <character> The levels of the actual <factor> |
| recall | <numeric> The recall |
| precision | <numeric> The precision |

Creating [<factor>](#)

Consider a classification problem with three classes: A, B, and C. The actual vector of [factor\(\)](#) values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

Definition

Let $\hat{\sigma} \in [0, 1]$ be the proportion of true negatives among the actual negatives. The specificity of the classifier is calculated as,

$$\hat{\sigma} = \frac{\#TN_k}{\#TN_k + \#FP_k}$$

Where:

- $\#TN_k$ is the number of true negatives, and
- $\#FP_k$ is the number of false positives.

See Also

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zeroonloss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroone.loss.factor()`

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data = iris,
  family = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
response <- predict(model, type = "response")

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 4) generate precision-recall
# data

# 4.1) calculate residual
# probability and store as matrix
response <- matrix(
  data = cbind(response, 1-response),
  nrow = length(actual)
)

# 4.2) generate precision-recall
# data
```

```

roc <- prROC(
  actual = actual,
  response = response
)

# 5) plot by species
plot(roc)

# 5.1) summarise
summary(roc)

# 6) provide custom
# thresholds
roc <- prROC(
  actual = actual,
  response = response,
  thresholds = seq(
    1,
    0,
    length.out = 20
  )
)

# 5) plot by species
plot(roc)

```

rae.numeric

Relative Absolute Error

Description

The `rae()`-function calculates the normalized **relative absolute error** between the predicted and observed `<numeric>` vectors. The `weighted.rae()` function computes the weighed relative absolute error.

Usage

```

## S3 method for class 'numeric'
rae(actual, predicted, ...)

## S3 method for class 'numeric'
weighted.rae(actual, predicted, w, ...)

## Generic S3 method
rae(
  actual,
  predicted,
  ...
)

```



```
## Generic S3 method
weighted.rae(
  actual,
  predicted,
  w,
  ...
)
```

Arguments

`actual` A <numeric>-vector of length n . The observed (continuous) response variable.

`predicted` A <numeric>-vector of length n . The estimated (continuous) response variable.

`...` Arguments passed into other methods.

`w` A <numeric>-vector of length n . The weight assigned to each observation in the data.

Value

A <numeric> vector of length 1.

Definition

The Relative Absolute Error (RAE) is calculated as:

$$\text{RAE} = \frac{\sum_{i=1}^n |y_i - v_i|}{\sum_{i=1}^n |y_i - \bar{y}|}$$

Where y_i are the actual values, v_i are the predicted values, and \bar{y} is the mean of the actual values.

See Also

Other Regression: `ccc.numeric()`, `huberloss.numeric()`, `mae.numeric()`, `mape.numeric()`, `mpe.numeric()`, `mse.numeric()`, `pinball.numeric()`, `rmse.numeric()`, `rmsle.numeric()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroonloss.factor()`

Examples

```

# 1) fit a linear
# regression
model <- lm(
  mpg ~ .,
  data = mtcars
)

# 1.1) define actual
# and predicted values
# to measure performance
actual <- mtcars$mpg
predicted <- fitted(model)

# 2) evaluate in-sample model
# performance using Relative Absolute Error (RAE)
cat(
  "Relative Absolute Error", rae(
    actual = actual,
    predicted = predicted,
  ),
  "Relative Absolute Error (weighted)", weighted.rae(
    actual = actual,
    predicted = predicted,
    w = mtcars$mpg/mean(mtcars$mpg)
  ),
  sep = "\n"
)

```

recall.factor

Recall

Description

A generic function for the **Recall**. Use `weighted.fdr()` for the weighted **Recall**.

Other names:

Sensitivity, True Positive Rate

Usage

```

## S3 method for class 'factor'
recall(actual, predicted, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
weighted.recall(actual, predicted, w, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'cmatrix'
recall(x, micro = NULL, na.rm = TRUE, ...)

```

```
## S3 method for class 'factor'
sensitivity(actual, predicted, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
weighted.sensitivity(actual, predicted, w, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'cmatrix'
sensitivity(x, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
tpr(actual, predicted, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
weighted.tpr(actual, predicted, w, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'cmatrix'
tpr(x, micro = NULL, na.rm = TRUE, ...)

## Generic S3 method
recall(
  ...,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
sensitivity(
  ...,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
tpr(
  ...,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
weighted.recall(
  ...,
  w,
  micro = NULL,
  na.rm = TRUE
)
```

```
## Generic S3 method
weighted.sensitivity(
  ...,
  w,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
weighted.tpr(
  ...,
  w,
  micro = NULL,
  na.rm = TRUE
)
```

Arguments

| | |
|-----------|--|
| actual | A vector of <factor> values of length n , and k levels. |
| predicted | A vector of <factor> values of length n , and k levels. |
| micro | A <logical> -value of length 1 (default: NULL). If TRUE it returns the micro average across all k classes, if FALSE it returns the macro average. |
| na.rm | A <logical> value of length 1 (default: TRUE). If TRUE , NA values are removed from the computation. This argument is only relevant when micro != NULL . When na.rm = TRUE , the computation corresponds to <code>sum(c(1, 2, NA), na.rm = TRUE) / length(na.omit(c(1, 2, NA)))</code> . When na.rm = FALSE , the computation corresponds to <code>sum(c(1, 2, NA), na.rm = TRUE) / length(c(1, 2, NA))</code> . |
| ... | Arguments passed into other methods |
| w | A <numeric> -vector of length n . NULL by default. |
| x | A confusion matrix created <code>cmatrix()</code> . |

Value

If **micro** is **NULL** (the default), a named **<numeric>**-vector of **length** k

If **micro** is **TRUE** or **FALSE**, a **<numeric>**-vector of **length** 1

Definition

Let $\hat{\rho} \in [0, 1]$ be the proportion of true positives among the actual positives. The recall of the classifier is calculated as,

$$\hat{\rho} = \frac{\#TP_k}{\#TP_k + \#FN_k}$$

Where:

- $\#TP_k$ is the number of true positives, and
- $\#FN_k$ is the number of false negatives.

Creating <factor>

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

See Also

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zeroonloss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroonloss.factor()`

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data    = iris,
  family  = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
predicted <- factor(
  as.numeric(
    predict(model, type = "response") > 0.5
  ),
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 4) evaluate class-wise performance
# using Recall

# 4.1) unweighted Recall
recall(
  actual    = actual,
  predicted = predicted
)

# 4.2) weighted Recall
weighted.recall(
  actual    = actual,
  predicted = predicted,
  w         = iris$Petal.Length/mean(iris$Petal.Length)
)
```

```

# 5) evaluate overall performance
# using micro-averaged Recall
cat(
  "Micro-averaged Recall", recall(
    actual = actual,
    predicted = predicted,
    micro = TRUE
  ),
  "Micro-averaged Recall (weighted)", weighted.recall(
    actual = actual,
    predicted = predicted,
    w = iris$Petal.Length/mean(iris$Petal.Length),
    micro = TRUE
  ),
  sep = "\n"
)

```

 rmse.numeric

Root Mean Squared Error

Description

The `rmse()`-function computes the **root mean squared error** between the observed and predicted `<numeric>` vectors. The `weighted.rmse()` function computes the weighted root mean squared error.

Usage

```

## S3 method for class 'numeric'
rmse(actual, predicted, ...)

## S3 method for class 'numeric'
weighted.rmse(actual, predicted, w, ...)

## Generic S3 method
rmse(
  actual,
  predicted,
  ...
)

weighted.rmse(
  actual,
  predicted,
  w,
  ...
)

```

Arguments

| | |
|-----------|---|
| actual | A <numeric>-vector of length n . The observed (continuous) response variable. |
| predicted | A <numeric>-vector of length n . The estimated (continuous) response variable. |
| ... | Arguments passed into other methods. |
| w | A <numeric>-vector of length n . The weight assigned to each observation in the data. |

Value

A <numeric> vector of length 1.

Definition

The metric is calculated as,

$$\sqrt{\frac{1}{n} \sum_i^n (y_i - v_i)^2}$$

Where y_i and v_i are the actual and predicted values respectively.

See Also

Other Regression: `ccc.numeric()`, `huberloss.numeric()`, `mae.numeric()`, `mape.numeric()`, `mpe.numeric()`, `mse.numeric()`, `pinball.numeric()`, `rae.numeric()`, `rmsle.numeric()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zerooneloss.factor()`

Examples

```
# 1) fit a linear
# regression
model <- lm(
  mpg ~ .,
  data = mtcars
)

# 1.1) define actual
# and predicted values
# to measure performance
actual <- mtcars$mpg
predicted <- fitted(model)
```



```

# 2) evaluate in-sample model
# performance using Root Mean Squared Error (RMSE)
cat(
  "Root Mean Squared Error", rmse(
    actual = actual,
    predicted = predicted,
  ),
  "Root Mean Squared Error (weighted)", weighted.rmse(
    actual = actual,
    predicted = predicted,
    w = mtcars$mpg/mean(mtcars$mpg)
  ),
  sep = "\n"
)

```

rmsle.numeric

Root Mean Squared Logarithmic Error

Description

The `rmsle()`-function computes the root mean squared logarithmic error between the observed and predicted `<numeric>` vectors. The `weighted.rmsle()` function computes the weighted root mean squared logarithmic error.

Usage

```

## S3 method for class 'numeric'
rmsle(actual, predicted, ...)

## S3 method for class 'numeric'
weighted.rmsle(actual, predicted, w, ...)

## Generic S3 method
rmsle(
  actual,
  predicted,
  ...
)

## Generic S3 method
weighted.rmsle(
  actual,
  predicted,
  w,
  ...
)

```

Arguments

| | |
|-----------|---|
| actual | A <numeric>-vector of length n . The observed (continuous) response variable. |
| predicted | A <numeric>-vector of length n . The estimated (continuous) response variable. |
| ... | Arguments passed into other methods. |
| w | A <numeric>-vector of length n . The weight assigned to each observation in the data. |

Value

A <numeric> vector of length 1.

Definition

The metric is calculated as,

$$\sqrt{\frac{1}{n} \sum_i^n (\log(1 + y_i) - \log(1 + v_i))^2}$$

Where y_i and v_i are the actual and predicted values respectively.

See Also

Other Regression: `ccc.numeric()`, `huberloss.numeric()`, `mae.numeric()`, `mape.numeric()`, `mpe.numeric()`, `mse.numeric()`, `pinball.numeric()`, `rae.numeric()`, `rmse.numeric()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zerooneloss.factor()`

Examples

```
# 1) fit a linear
# regression
model <- lm(
  mpg ~ .,
  data = mtcars
)

# 1.1) define actual
# and predicted values
# to measure performance
actual <- mtcars$mpg
predicted <- fitted(model)
```

```

# 2) evaluate in-sample model
# performance using Root Mean Squared Logarithmic Error (RMSLE)
cat(
  "Root Mean Squared Logarithmic Error", rmsle(
    actual = actual,
    predicted = predicted,
  ),
  "Root Mean Squared Logarithmic Error (weighted)", weighted.rmsle(
    actual = actual,
    predicted = predicted,
    w = mtcars$mpg/mean(mtcars$mpg)
  ),
  sep = "\n"
)

```

roc.auc.matrix

Area under the Receiver Operator Characteristics Curve

Description

A generic function for the area under the Receiver Operator Characteristics Curve. Use [weighted.roc.auc\(\)](#) for the weighted area under the Receiver Operator Characteristics Curve.

Usage

```

## S3 method for class 'matrix'
roc.auc(actual, response, micro = NULL, method = 0L, ...)

## S3 method for class 'matrix'
weighted.roc.auc(actual, response, w, micro = NULL, method = 0L, ...)

## Generic S3 method
roc.auc(
  actual,
  response,
  micro = NULL,
  method = 0,
  ...
)

## Generic S3 method
weighted.roc.auc(
  actual,
  response,
  w,
  micro = NULL,
  method = 0,

```

...
)

Arguments

| | |
|----------|---|
| actual | A vector of <factor> values of length n , and k levels. |
| response | A $n \times k$ <numeric>-matrix. The estimated response probabilities for each class k . |
| micro | A <logical>-value of length 1 (default: <code>NULL</code>). If <code>TRUE</code> it returns the micro average across all k classes, if <code>FALSE</code> it returns the macro average. |
| method | A <numeric> value (default: 0). Defines the underlying method of calculating the area under the curve. If 0 it is calculated using the trapezoid-method, if 1 it is calculated using the step-method. |
| ... | Arguments passed into other methods. |
| w | A <numeric>-vector of length n . <code>NULL</code> by default. |

Value

A <numeric> vector of length 1

Definition

Trapezoidal rule

The **trapezoidal rule** approximates the integral of a function $f(x)$ between $x = a$ and $x = b$ using trapezoids formed between consecutive points. If we have points x_0, x_1, \dots, x_n (with $a = x_0 < x_1 < \dots < x_n = b$) and corresponding function values $f(x_0), f(x_1), \dots, f(x_n)$, the area under the curve A_T is approximated by:

$$A_T \approx \sum_{k=1}^n \frac{f(x_{k-1}) + f(x_k)}{2} [x_k - x_{k-1}].$$

Step-function method

The **step-function (rectangular) method** uses the value of the function at one endpoint of each subinterval to form rectangles. With the same partition x_0, x_1, \dots, x_n , the rectangular approximation A_S can be written as:

$$A_S \approx \sum_{k=1}^n f(x_{k-1}) [x_k - x_{k-1}].$$

See Also

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `specificity.factor()`, `zeroonloss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zerooneloss.factor()`

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `specificity.factor()`, `zerooneloss.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zerooneloss.factor()`

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data = iris,
  family = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
response <- predict(model, type = "response")

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)
```

```

# 4) generate receiver operator characteristics
# data

# 4.1) calculate residual
# probability and store as matrix
response <- matrix(
  data = cbind(response, 1-response),
  nrow = length(actual)
)

# 4.2) calculate class-wise
# area under the curve
roc.auc(
  actual = actual,
  response = response
)

# 4.3) calculate class-wise
# weighted area under the curve
weighted.roc.auc(
  actual = actual,
  response = response,
  w = iris$Petal.Length/mean(iris$Petal.Length)
)

# 5) evaluate overall area under
# the curve
cat(
  "Micro-averaged area under the ROC curve", roc.auc(
    actual = actual,
    response = response,
    micro = TRUE
  ),
  "Micro-averaged area under the ROC curve (weighted)", weighted.roc.auc(
    actual = actual,
    response = response,
    w = iris$Petal.Length/mean(iris$Petal.Length),
    micro = TRUE
  ),
  sep = "\n"
)

```

ROC.factor

Receiver Operator Characteristics

Description

The `ROC()`-function computes the `tpr()` and `fpr()` at thresholds provided by the `response`- or `thresholds`-vector. The function constructs a `data.frame()` grouped by k -classes where each class is treated as a binary classification problem.

Usage

```
## S3 method for class 'factor'
ROC(actual, response, thresholds = NULL, presorted = FALSE, ...)

## S3 method for class 'factor'
weighted.ROC(actual, response, w, thresholds = NULL, presorted = FALSE, ...)

## Generic S3 method
ROC(
  actual,
  response,
  thresholds = NULL,
  presorted = FALSE,
  ...
)

## Generic S3 method
weighted.ROC(
  actual,
  response,
  w,
  thresholds = NULL,
  presorted = FALSE,
  ...
)
```

Arguments

| | |
|------------|---|
| actual | A vector of <code><factor></code> values of <code>length</code> n , and k levels. |
| response | A $n \times k$ <code><numeric></code> -matrix. The estimated response probabilities for each class k . |
| thresholds | An optional <code><numeric></code> vector of <code>length</code> n (default: <code>NULL</code>). |
| presorted | A <code><logical></code> -value <code>length</code> 1 (default: <code>FALSE</code>). If <code>TRUE</code> the input will not be sorted by threshold. |
| ... | Arguments passed into other methods. |
| w | A <code><numeric></code> -vector of <code>length</code> n . <code>NULL</code> by default. |

Value

A `data.frame` on the following form,

| | |
|-----------|---|
| threshold | <code><numeric></code> Thresholds used to determine <code>tpr()</code> and <code>fpr()</code> |
| level | <code><character></code> The level of the actual <code><factor></code> |
| label | <code><character></code> The levels of the actual <code><factor></code> |
| fpr | <code><numeric></code> The false positive rate |
| tpr | <code><numeric></code> The true positive rate |

Creating <factor>

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

Definition

Let $\hat{\sigma} \in [0, 1]$ be the proportion of true negatives among the actual negatives. The specificity of the classifier is calculated as,

$$\hat{\sigma} = \frac{\#TN_k}{\#TN_k + \#FP_k}$$

Where:

- $\#TN_k$ is the number of true negatives, and
- $\#FP_k$ is the number of false positives.

See Also

Other Classification: `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`, `zeroone.loss.factor()`

Other Supervised Learning: `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroone.loss.factor()`

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data = iris,
  family = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
response <- predict(model, type = "response")

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 4) generate receiver
# operator characteristics

# 4.1) calculate residual
# probability and store as matrix
```

```

response <- matrix(
  data = cbind(response, 1-response),
  nrow = length(actual)
)

# 4.2) construct
# data.frame
roc <- ROC(
  actual = actual,
  response = response
)

# 5) plot by species
plot(roc)

# 5.1) summarise
summary(roc)

# 6) provide custom
# thresholds
roc <- ROC(
  actual = actual,
  response = response,
  thresholds = seq(
    1,
    0,
    length.out = 20
  )
)

# 5) plot by species
plot(roc)

```

rrmse.numeric

Relative Root Mean Squared Error

Description

The `rrmse()`-function computes the **Relative Root Mean Squared Error** between the observed and predicted `<numeric>` vectors. The `weighted.rrmse()` function computes the weighted Relative Root Mean Squared Error.

Usage

```

## S3 method for class 'numeric'
rrmse(actual, predicted, normalization = 1L, ...)

## S3 method for class 'numeric'
weighted.rrmse(actual, predicted, w, normalization = 1L, ...)

```

```
## Generic S3 method
rrmse(
  actual,
  predicted,
  normalization = 1,
  ...
)

## Generic S3 method
weighted.rrmse(
  actual,
  predicted,
  w,
  normalization = 1,
  ...
)
```

Arguments

| | |
|---------------|--|
| actual | A <numeric> -vector of length n . The observed (continuous) response variable. |
| predicted | A <numeric> -vector of length n . The estimated (continuous) response variable. |
| normalization | A <numeric> -value of length 1 (default: 1). 0: mean-normalization , 1: range-normalization , 2: IQR-normalization . |
| ... | Arguments passed into other methods. |
| w | A <numeric> -vector of length n . The weight assigned to each observation in the data. |

Value

A [<numeric>](#) vector of [length](#) 1.

Definition

The metric is calculated as,

$$\frac{RMSE}{\gamma}$$

Where γ is the normalization factor.

See Also

Other Regression: [ccc.numeric\(\)](#), [huberloss.numeric\(\)](#), [mae.numeric\(\)](#), [mape.numeric\(\)](#), [mpe.numeric\(\)](#), [mse.numeric\(\)](#), [pinball.numeric\(\)](#), [rae.numeric\(\)](#), [rmse.numeric\(\)](#), [rmsle.numeric\(\)](#), [rrse.numeric\(\)](#), [rsq.numeric\(\)](#), [smape.numeric\(\)](#)

Other Supervised Learning: [ROC.factor\(\)](#), [accuracy.factor\(\)](#), [baccuracy.factor\(\)](#), [ccc.numeric\(\)](#), [ckappa.factor\(\)](#), [cmatrix.factor\(\)](#), [dor.factor\(\)](#), [entropy.matrix\(\)](#), [fbeta.factor\(\)](#),

```
fdr.factor(), fer.factor(), fpr.factor(), huberloss.numeric(), jaccard.factor(), logloss.factor(),
mae.numeric(), mape.numeric(), mcc.factor(), mpe.numeric(), mse.numeric(), nlr.factor(),
npv.factor(), pinball.numeric(), plr.factor(), pr.auc.matrix(), prROC.factor(), precision.factor(),
rae.numeric(), recall.factor(), rmse.numeric(), rmsle.numeric(), roc.auc.matrix(),
rrse.numeric(), rsq.numeric(), smape.numeric(), specificity.factor(), zeroone.loss.factor()
```

Examples

```
# 1) fit a linear
# regression
model <- lm(
  mpg ~ .,
  data = mtcars
)

# 1.1) define actual
# and predicted values
# to measure performance
actual <- mtcars$mpg
predicted <- fitted(model)

# 2) evaluate in-sample model
# performance using Relative Root Mean Squared Error (RRMSE)
cat(
  "IQR Relative Root Mean Squared Error", rmse(
    actual = actual,
    predicted = predicted,
    normalization = 2
  ),
  "IQR Relative Root Mean Squared Error (weighted)", weighted.rmse(
    actual = actual,
    predicted = predicted,
    w = mtcars$mpg/mean(mtcars$mpg),
    normalization = 2
  ),
  sep = "\n"
)
```

 rrse.numeric

Root Relative Squared Error

Description

The `rrse()`-function calculates the **root relative squared error** between the predicted and observed `<numeric>` vectors. The `weighted.rrse()` function computes the weighed root relative squared error.

Usage

```
## S3 method for class 'numeric'
rrse(actual, predicted, ...)

## S3 method for class 'numeric'
weighted.rrse(actual, predicted, w, ...)

## Generic S3 method
rrse(
  actual,
  predicted,
  ...
)

## Generic S3 method
weighted.rrse(
  actual,
  predicted,
  w,
  ...
)
```

Arguments

| | |
|-----------|---|
| actual | A <numeric>-vector of length n . The observed (continuous) response variable. |
| predicted | A <numeric>-vector of length n . The estimated (continuous) response variable. |
| ... | Arguments passed into other methods. |
| w | A <numeric>-vector of length n . The weight assigned to each observation in the data. |

Value

A <numeric> vector of length 1.

Definition

The metric is calculated as,

$$\text{RRSE} = \sqrt{\frac{\sum_{i=1}^n (y_i - v_i)^2}{\sum_{i=1}^n (y_i - \bar{y})^2}}$$

Where y_i are the actual values, v_i are the predicted values, and \bar{y} is the mean of the actual values.

See Also

Other Regression: `ccc.numeric()`, `huberloss.numeric()`, `mae.numeric()`, `mape.numeric()`, `mpe.numeric()`, `mse.numeric()`, `pinball.numeric()`, `rae.numeric()`, `rmse.numeric()`, `rmsle.numeric()`, `rrmse.numeric()`, `rsq.numeric()`, `smape.numeric()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rsq.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroonloss.factor()`

Examples

```
# 1) fit a linear
# regression
model <- lm(
  mpg ~ .,
  data = mtcars
)

# 1.1) define actual
# and predicted values
# to measure performance
actual <- mtcars$mpg
predicted <- fitted(model)

# 2) evaluate in-sample model
# performance using Relative Root Squared Error (RRSE)
cat(
  "Relative Root Squared Error", rrse(
    actual = actual,
    predicted = predicted,
  ),
  "Relative Root Squared Error (weighted)", weighted.rrse(
    actual = actual,
    predicted = predicted,
    w = mtcars$mpg/mean(mtcars$mpg)
  ),
  sep = "\n"
)
```

rsq.numeric

 R^2

Description

A generic function for the R^2 . The unadjusted R^2 is returned by default. Use `weighted.rsq()` for the weighted R^2 .

Usage

```
## S3 method for class 'numeric'
rsq(actual, predicted, k = 0, ...)

## S3 method for class 'numeric'
weighted.rsq(actual, predicted, w, k = 0, ...)

## Generic S3 method
rsq(
  ...,
  k = 0
)

## Generic S3 method
weighted.rsq(
  ...,
  w,
  k = 0
)
```

Arguments

actual A <numeric>-vector of **length** n . The observed (continuous) response variable.

predicted A <numeric>-vector of **length** n . The estimated (continuous) response variable.

k A <numeric>-vector of **length** 1 (default: 0). For adjusted R^2 set $k = \kappa - 1$, where κ is the number of parameters.

... Arguments passed into other methods.

w A <numeric>-vector of **length** n . The weight assigned to each observation in the data.

Value

A <numeric> vector of **length** 1.

Definition

Let $R^2 \in [-\infty, 1]$ be the explained variation. The R^2 is calculated as,

$$R^2 = 1 - \frac{\sum (y_i - \hat{y}_i)^2}{\sum (y_i - \bar{y})^2} \frac{n - 1}{n - (k + 1)}$$

Where:

- n is the number of observations
- k is the number of features
- y is the actual values
- \hat{y}_i is the predicted values

- $\sum (y_i - \hat{y}_i)^2$ is the sum of squared errors and,
- $\sum (y_i - \bar{y})^2$ is total sum of squared errors

See Also

Other Regression: `ccc.numeric()`, `huberloss.numeric()`, `mae.numeric()`, `mape.numeric()`, `mpe.numeric()`, `mse.numeric()`, `pinball.numeric()`, `rae.numeric()`, `rmse.numeric()`, `rmsle.numeric()`, `rrmse.numeric()`, `rrse.numeric()`, `smape.numeric()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`, `npv.factor()`, `pinball.numeric()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `rae.numeric()`, `recall.factor()`, `rmse.numeric()`, `rmsle.numeric()`, `roc.auc.matrix()`, `rrmse.numeric()`, `rrse.numeric()`, `smape.numeric()`, `specificity.factor()`, `zeroonloss.factor()`

Examples

```
# 1) fit a linear
# regression
model <- lm(
  mpg ~ .,
  data = mtcars
)

# 1.1) define actual
# and predicted values
# to measure in-sample performance
actual <- mtcars$mpg
predicted <- fitted(model)

# 2) calculate performance
# using R squared adjusted and
# unadjusted for features
cat(
  "Rsquared", rsq(
    actual = actual,
    predicted = fitted(model)
  ),
  "Rsquared (Adjusted)", rsq(
    actual = actual,
    predicted = fitted(model),
    k = ncol(model.matrix(model)) - 1
  ),
  sep = "\n"
)
```

`smape.numeric`*Symmetric Mean Absolute Percentage Error*

Description

The `smape()`-function computes the **symmetric mean absolute percentage error** between the observed and predicted `<numeric>` vectors. The `weighted.smape()` function computes the weighted symmetric mean absolute percentage error.

Usage

```
## S3 method for class 'numeric'
smape(actual, predicted, ...)

## S3 method for class 'numeric'
weighted.smape(actual, predicted, w, ...)

## Generic S3 method
smape(
  actual,
  predicted,
  ...
)

## Generic S3 method
weighted.smape(
  actual,
  predicted,
  w,
  ...
)
```

Arguments

| | |
|------------------------|--|
| <code>actual</code> | A <code><numeric></code> -vector of length n . The observed (continuous) response variable. |
| <code>predicted</code> | A <code><numeric></code> -vector of length n . The estimated (continuous) response variable. |
| <code>...</code> | Arguments passed into other methods. |
| <code>w</code> | A <code><numeric></code> -vector of length n . The weight assigned to each observation in the data. |

Value

A `<numeric>` vector of **length** 1.

Definition

The metric is calculated as follows,

$$\sum_i^n \frac{1}{n} \frac{|y_i - v_i|}{\frac{|y_i| + |v_i|}{2}}$$

where y_i and v_i is the actual and predicted values respectively.

See Also

Other Regression: [ccc.numeric\(\)](#), [huberloss.numeric\(\)](#), [mae.numeric\(\)](#), [mape.numeric\(\)](#), [mpe.numeric\(\)](#), [mse.numeric\(\)](#), [pinball.numeric\(\)](#), [rae.numeric\(\)](#), [rmse.numeric\(\)](#), [rmsle.numeric\(\)](#), [rrmse.numeric\(\)](#), [rrse.numeric\(\)](#), [rsq.numeric\(\)](#)

Other Supervised Learning: [ROC.factor\(\)](#), [accuracy.factor\(\)](#), [baccuracy.factor\(\)](#), [ccc.numeric\(\)](#), [ckappa.factor\(\)](#), [cmatrix.factor\(\)](#), [dor.factor\(\)](#), [entropy.matrix\(\)](#), [fbeta.factor\(\)](#), [fdr.factor\(\)](#), [fer.factor\(\)](#), [fpr.factor\(\)](#), [huberloss.numeric\(\)](#), [jaccard.factor\(\)](#), [logloss.factor\(\)](#), [mae.numeric\(\)](#), [mape.numeric\(\)](#), [mcc.factor\(\)](#), [mpe.numeric\(\)](#), [mse.numeric\(\)](#), [nlr.factor\(\)](#), [npv.factor\(\)](#), [pinball.numeric\(\)](#), [plr.factor\(\)](#), [pr.auc.matrix\(\)](#), [prROC.factor\(\)](#), [precision.factor\(\)](#), [rae.numeric\(\)](#), [recall.factor\(\)](#), [rmse.numeric\(\)](#), [rmsle.numeric\(\)](#), [roc.auc.matrix\(\)](#), [rrmse.numeric\(\)](#), [rrse.numeric\(\)](#), [rsq.numeric\(\)](#), [specificity.factor\(\)](#), [zeroonloss.factor\(\)](#)

Examples

```
# 1) fit a linear
# regression
model <- lm(
  mpg ~ .,
  data = mtcars
)

# 1.1) define actual
# and predicted values
# to measure performance
actual <- mtcars$mpg
predicted <- fitted(model)

# 2) evaluate in-sample model
# performance using Symmetric Mean Absolute Percentage Error (MAPE)
cat(
  "Symmetric Mean Absolute Percentage Error", mape(
    actual = actual,
    predicted = predicted,
  ),
  "Symmetric Mean Absolute Percentage Error (weighted)", weighted.mape(
    actual = actual,
    predicted = predicted,
    w = mtcars$mpg/mean(mtcars$mpg)
  ),
  sep = "\n"
)
```

| | |
|--------------------|--------------------|
| specificity.factor | <i>Specificity</i> |
|--------------------|--------------------|

Description

A generic function for the **Specificity**. Use `weighted.specificity()` for the weighted **Specificity**.

Other names:

True Negative Rate, Selectivity

Usage

```
## S3 method for class 'factor'
specificity(actual, predicted, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
weighted.specificity(actual, predicted, w, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'cmatrix'
specificity(x, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
tnr(actual, predicted, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
weighted.tnr(actual, predicted, w, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'cmatrix'
tnr(x, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
selectivity(actual, predicted, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'factor'
weighted.selectivity(actual, predicted, w, micro = NULL, na.rm = TRUE, ...)

## S3 method for class 'cmatrix'
selectivity(x, micro = NULL, na.rm = TRUE, ...)

## Generic S3 method
specificity(
  ...,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
```

```

tnr(
  ...,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
selectivity(
  ...,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
weighted.specificity(
  ...,
  w,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
weighted.tnr(
  ...,
  w,
  micro = NULL,
  na.rm = TRUE
)

## Generic S3 method
weighted.selectivity(
  ...,
  w,
  micro = NULL,
  na.rm = TRUE
)

```

Arguments

| | |
|-----------|---|
| actual | A vector of <code><factor></code> values of length n , and k levels. |
| predicted | A vector of <code><factor></code> values of length n , and k levels. |
| micro | A <code><logical></code> -value of length 1 (default: <code>NULL</code>). If <code>TRUE</code> it returns the micro average across all k classes, if <code>FALSE</code> it returns the macro average. |
| na.rm | A <code><logical></code> value of length 1 (default: <code>TRUE</code>). If <code>TRUE</code> , <code>NA</code> values are removed from the computation. This argument is only relevant when <code>micro != NULL</code> . When <code>na.rm = TRUE</code> , the computation corresponds to <code>sum(c(1, 2, NA), na.rm = TRUE) / length(na.omit(c(1, 2, NA)))</code> . When <code>na.rm = FALSE</code> , the |

computation corresponds to `sum(c(1, 2, NA), na.rm = TRUE) / length(c(1, 2, NA))`.

... Arguments passed into other methods

w A <numeric>-vector of length *n*. `NULL` by default.

x A confusion matrix created `cmatrix()`.

Value

If `micro` is `NULL` (the default), a named <numeric>-vector of length *k*

If `micro` is `TRUE` or `FALSE`, a <numeric>-vector of length 1

Creating <factor>

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

Definition

Let $\hat{\sigma} \in [0, 1]$ be the proportion of true negatives among the actual negatives. The specificity of the classifier is calculated as,

$$\hat{\sigma} = \frac{\#TN_k}{\#TN_k + \#FP_k}$$

Where:

- $\#TN_k$ is the number of true negatives, and
- $\#FP_k$ is the number of false positives.

See Also

Other Classification: [ROC.factor\(\)](#), [accuracy.factor\(\)](#), [baccuracy.factor\(\)](#), [ckappa.factor\(\)](#), [cmatrix.factor\(\)](#), [dor.factor\(\)](#), [entropy.matrix\(\)](#), [fbeta.factor\(\)](#), [fdr.factor\(\)](#), [fer.factor\(\)](#), [fmi.factor\(\)](#), [fpr.factor\(\)](#), [jaccard.factor\(\)](#), [logloss.factor\(\)](#), [mcc.factor\(\)](#), [nlr.factor\(\)](#), [npv.factor\(\)](#), [plr.factor\(\)](#), [pr.auc.matrix\(\)](#), [prROC.factor\(\)](#), [precision.factor\(\)](#), [recall.factor\(\)](#), [roc.auc.matrix\(\)](#), [zerooneloss.factor\(\)](#)

Other Supervised Learning: [ROC.factor\(\)](#), [accuracy.factor\(\)](#), [baccuracy.factor\(\)](#), [ccc.numeric\(\)](#), [ckappa.factor\(\)](#), [cmatrix.factor\(\)](#), [dor.factor\(\)](#), [entropy.matrix\(\)](#), [fbeta.factor\(\)](#), [fdr.factor\(\)](#), [fer.factor\(\)](#), [fpr.factor\(\)](#), [huberloss.numeric\(\)](#), [jaccard.factor\(\)](#), [logloss.factor\(\)](#), [mae.numeric\(\)](#), [mape.numeric\(\)](#), [mcc.factor\(\)](#), [mpe.numeric\(\)](#), [mse.numeric\(\)](#), [nlr.factor\(\)](#), [npv.factor\(\)](#), [pinball.numeric\(\)](#), [plr.factor\(\)](#), [pr.auc.matrix\(\)](#), [prROC.factor\(\)](#), [precision.factor\(\)](#), [rae.numeric\(\)](#), [recall.factor\(\)](#), [rmse.numeric\(\)](#), [rmsle.numeric\(\)](#), [roc.auc.matrix\(\)](#), [rrmse.numeric\(\)](#), [rrse.numeric\(\)](#), [rsq.numeric\(\)](#), [smape.numeric\(\)](#), [zerooneloss.factor\(\)](#)

Examples

```
# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data = iris,
  family = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
predicted <- factor(
  as.numeric(
```

```
    predict(model, type = "response") > 0.5
  ),
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 4) evaluate class-wise performance
# using Specificity

# 4.1) unweighted Specificity
specificity(
  actual = actual,
  predicted = predicted
)

# 4.2) weighted Specificity
weighted.specificity(
  actual = actual,
  predicted = predicted,
  w = iris$Petal.Length/mean(iris$Petal.Length)
)

# 5) evaluate overall performance
# using micro-averaged Specificity
cat(
  "Micro-averaged Specificity", specificity(
    actual = actual,
    predicted = predicted,
    micro = TRUE
  ),
  "Micro-averaged Specificity (weighted)", weighted.specificity(
    actual = actual,
    predicted = predicted,
    w = iris$Petal.Length/mean(iris$Petal.Length),
    micro = TRUE
  ),
  sep = "\n"
)
```

Description

This dataset contains measurements of various chemical properties of white wines along with their quality ratings and a quality classification. The dataset was obtained from the UCI Machine Learning Repository.

Usage

```
data(wine_quality)
```

Format

A list with two components:

features A data frame with 11 chemical property variables.

target A list with two elements: regression (wine quality scores) and class (quality classification).

Details

The data is provided as a list with two components:

features A data frame containing the chemical properties of the wines. The variables include:

fixed_acidity Fixed acidity (g/L).

volatile_acidity Volatile acidity (g/L), mainly due to acetic acid.

citric_acid Citric acid (g/L).

residual_sugar Residual sugar (g/L).

chlorides Chloride concentration (g/L).

free_sulfur_dioxide Free sulfur dioxide (mg/L).

total_sulfur_dioxide Total sulfur dioxide (mg/L).

density Density of the wine (g/cm³).

pH pH value of the wine.

sulphates Sulphates (g/L).

alcohol Alcohol content (% by volume).

target A list containing two elements:

regression A numeric vector representing the wine quality scores (used as the regression target).

class A factor with levels "High Quality", "Medium Quality", and "Low Quality", where classification is determined as follows:

High Quality quality ≥ 7 .

Low Quality quality ≤ 4 .

Medium Quality for all other quality scores.

Source

<https://archive.ics.uci.edu/dataset/186/wine+quality>

 zerooneLoss.factor *Zero-One Loss*

Description

The `zerooneLoss()`-function computes the **zero-one Loss**, a classification loss function that calculates the proportion of misclassified instances between two vectors of predicted and observed `factor()` values. The `weighted.zerooneLoss()` function computes the weighted zero-one loss.

Usage

```
## S3 method for class 'factor'
zerooneLoss(actual, predicted, ...)

## S3 method for class 'factor'
weighted.zerooneLoss(actual, predicted, w, ...)

## S3 method for class 'cmatrix'
zerooneLoss(x, ...)

## Generic S3 method
zerooneLoss(...)

## Generic S3 method
weighted.zerooneLoss(
  ...,
  w
)
```

Arguments

| | |
|-----------|---|
| actual | A vector of <code><factor></code> with length n , and k levels |
| predicted | A vector of <code><factor></code> with length n , and k levels |
| ... | Arguments passed into other methods |
| w | A <code><numeric></code> -vector of length n . <code>NULL</code> by default |
| x | A confusion matrix created <code>cmatrix()</code> |

Value

A `<numeric>`-vector of length 1

Definition

The metric is calculated as follows,

$$\frac{\#FP + \#FN}{\#TP + \#TN + \#FP + \#FN}$$

Where $\#TP$, $\#TN$, $\#FP$, and $\#FN$ represent the true positives, true negatives, false positives, and false negatives, respectively.

Creating `<factor>`

Consider a classification problem with three classes: A, B, and C. The actual vector of `factor()` values is defined as follows:

```
## set seed
set.seed(1903)

## actual
factor(
  x = sample(x = 1:3, size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] B A B B A C B C C A
#> Levels: A B C
```

Here, the values 1, 2, and 3 are mapped to A, B, and C, respectively. Now, suppose your model does not predict any B's. The predicted vector of `factor()` values would be defined as follows:

```
## set seed
set.seed(1903)

## predicted
factor(
  x = sample(x = c(1, 3), size = 10, replace = TRUE),
  levels = c(1, 2, 3),
  labels = c("A", "B", "C")
)
#> [1] C A C C C C C C A C
#> Levels: A B C
```

In both cases, $k = 3$, determined indirectly by the `levels` argument.

See Also

Other Classification: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fmi.factor()`, `fpr.factor()`, `jaccard.factor()`, `logloss.factor()`, `mcc.factor()`, `nlr.factor()`, `npv.factor()`, `plr.factor()`, `pr.auc.matrix()`, `prROC.factor()`, `precision.factor()`, `recall.factor()`, `roc.auc.matrix()`, `specificity.factor()`

Other Supervised Learning: `ROC.factor()`, `accuracy.factor()`, `baccuracy.factor()`, `ccc.numeric()`, `ckappa.factor()`, `cmatrix.factor()`, `dor.factor()`, `entropy.matrix()`, `fbeta.factor()`, `fdr.factor()`, `fer.factor()`, `fpr.factor()`, `huberloss.numeric()`, `jaccard.factor()`, `logloss.factor()`, `mae.numeric()`, `mape.numeric()`, `mcc.factor()`, `mpe.numeric()`, `mse.numeric()`, `nlr.factor()`,

```

npv.factor(), pinball.numeric(), plr.factor(), pr.auc.matrix(), prROC.factor(), precision.factor(),
rae.numeric(), recall.factor(), rmse.numeric(), rmsle.numeric(), roc.auc.matrix(),
rrmse.numeric(), rrse.numeric(), rsq.numeric(), smape.numeric(), specificity.factor()

```

Examples

```

# 1) recode Iris
# to binary classification
# problem
iris$species_num <- as.numeric(
  iris$Species == "virginica"
)

# 2) fit the logistic
# regression
model <- glm(
  formula = species_num ~ Sepal.Length + Sepal.Width,
  data = iris,
  family = binomial(
    link = "logit"
  )
)

# 3) generate predicted
# classes
predicted <- factor(
  as.numeric(
    predict(model, type = "response") > 0.5
  ),
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 3.1) generate actual
# classes
actual <- factor(
  x = iris$species_num,
  levels = c(1,0),
  labels = c("Virginica", "Others")
)

# 4) evaluate model
# performance using Zero-One Loss
cat(
  "Zero-One Loss", zeroone.loss(
    actual = actual,
    predicted = predicted
  ),
  "Zero-One Loss (weigthed)", weighted.zeroone.loss(
    actual = actual,
    predicted = predicted,
    w = iris$Petal.Length/mean(iris$Petal.Length)
  ),

```

```
    sep = "\n"  
  )
```

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