# Package 'HPLB'

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Title High-Probability Lower Bounds for the Total Variance Distance
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Author Loris Michel, Jeffrey Naef
Maintainer Loris Michel <michel@stat.math.ethz.ch></michel@stat.math.ethz.ch>
<b>Description</b> An implementation of high-probability lower bounds for the total variance distance as introduced in Michel & Naef & Meinshausen (2020) <arxiv:2005.06006>. An estimated lower-bound (with high-probability) on the total variation distance between two probability distributions from which samples are observed can be obtained with the function HPLB.</arxiv:2005.06006>
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boundingOperation	<b>Bounding Operation</b>
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## **Description**

**Bounding Operation** 

## Usage

```
boundingOperation(v, left, right, m, n)
```

## **Arguments**

V	a a numeric value giving an ordering permutation of 1 to m+n.
left	a numeric value giving the number of witnesses left.
right	a numeric value giving the number of witnesses right.
m	a numeric value, the number of observations left.
n	a numeric value, the number of observations right.

#### Value

a cumulative counting function represented as a numeric vector.

empiricalBF	Empirical Bounding Functions

## Description

**Empirical Bounding Functions** 

## Usage

```
empiricalBF(tv.seq, nsim = 1000, m = 100, n = 100, alpha = 0.05)
```

## **Arguments**

```
tv.seq a vector of total variation values between 0 and 1.

nsim a numeric value giving the number of repetitions.

m a numeric value, the number of observations left.

n a numeric value, the number of observations right.

alpha a numeric value giving the type-I error level.
```

## Value

a list of empirical bounding functions indexed by the tv.seq (in the respective order).

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**HPLB** 

High Probability Lower Bounds (HPLB) for the Total Variation Distance (TV) Based on Finite Samples

#### **Description**

Implementations of different HPLBs for TV as described in (Michel et al., 2020).

#### Usage

```
HPLB(
    t,
    rho,
    s = 0.5,
    estimator.type = "adapt",
    alpha = 0.05,
    tv.seq = seq(from = 0, to = 1, by = 1/length(t)),
    custom.bounding.seq = NULL,
    direction = rep("left", length(s)),
    cutoff = 0.5,
    verbose.plot = FALSE,
    seed = 0,
    ...
)
```

## **Arguments**

t a numeric vector value corresponding to a natural ordering of the observations.

For a two-sample test 0-1 numeric values values should be provided.

rho a numeric vector value providing an ordering. This could be a binary classifier, a regressor, a witness function from a MMD kernel or anything else that would

witness a distributional difference.

s a numeric vector value giving split points on t.

estimator.type a character value indicating which estimator to use. One option out of:

- adapt:adaptive binary classification estimator (asymptotic bounding function)
- · bayes:binary classification estimator
- bayes\_finite\_sample:binary classification finite sample estimator
- adapt\_empirical:adaptive binary classification estimator (simulation-based bounding function)
- adapt\_custom:adaptive binary classificatrion estimator (user-defined bounding function)
- adapt\_dwit:adaptive binary classificatrion estimator (for distributional witnesses estimation)

alpha a numeric value giving the overall type-I error control level.

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tv.seq a sequence of values between 0 and 1 used as the grid search for the total variation distance in case of tv-search.

custom.bounding.seq a list of bounding functions respecting the order of tv.seq used in case of estimator.type "custom-tv-search".

direction a character vector value made of "left" or "right" giving which distribution witness count to estimate (t<=s or t>s?).

cutoff a numeric value. This is the cutoff used if bayes estimators are used. The theory suggests to use 1/2 but this can be changed.

verbose.plot a boolean value for additional plots.

seed an integer value. The seed for reproducibility.

... additional parameters for the function empiricalBF.

#### Value

a list containing the relevant lower bounds estimates. For the total variation distance the relevant entry is tvhat.

#### Author(s)

Loris Michel, Jeffrey Naef

#### References

L. Michel, J. Naef and N. Meinshausen (2020). High-Probability Lower Bounds for the Total Variation Distance

## **Examples**

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```
# estimated HPLB
tvhat <- HPLB(t = t, rho = bayesRate(x), estimator.type = "bayes")</pre>
TotalVarDist(e1 = Norm(2,1), e2 = Norm(0,1))
## Example 2: optimal mixture detection (adapt estimator), Gaussian mean-shift example
n <- 100
mean.shift <- 2
t.train <- runif(n, 0 ,1)</pre>
x.train <- ifelse(t.train>0.5, stats::rnorm(n, mean.shift), stats::rnorm(n))
rf <- ranger::ranger(t~x, data.frame(t=t.train,x=x.train))</pre>
n <- 100
t.test <- runif(n, 0 ,1)</pre>
x.test <- ifelse(t.test>0.5, stats::rnorm(n, mean.shift), stats::rnorm(n))
rho <- predict(rf, data.frame(t=t.test,x=x.test))$predictions</pre>
## out-of-sample
tv.oos <- HPLB(t = t.test, rho = rho, s = seq(0.1,0.9,0.1), estimator.type = "adapt")
## total variation values
tv <- c()
for (s in seq(0.1,0.9,0.1)) {
 if (s<=0.5) {
   D.left <- Norm(0,1)
 } else {
   D.left <- UnivarMixingDistribution(Dlist = list(Norm(0,1),Norm(mean.shift,1)),</pre>
               mixCoeff = c(ifelse(s <= 0.5, 1, 0.5/s), ifelse(s <= 0.5, 0, (s - 0.5)/s)))
 if (s < 0.5) {
   D.right <- UnivarMixingDistribution(Dlist = list(Norm(0,1),Norm(mean.shift,1)),</pre>
           mixCoeff = c(ifelse(s <= 0.5, (0.5-s)/(1-s), 0), ifelse(s <= 0.5, (0.5/(1-s)), 1)))
 } else {
   D.right <- Norm(mean.shift,1)</pre>
tv <- c(tv, TotalVarDist(e1 = D.left, e2 = D.right))</pre>
}
## plot
oldpar <- par(no.readonly =TRUE)</pre>
par(mfrow=c(2,1))
plot(t.test,x.test,pch=19,xlab="t",ylab="x")
plot(seq(0.1,0.9,0.1), tv.oos$tvhat,type="l",ylim=c(0,1),xlab="t", ylab="TV")
lines(seq(0.1, 0.9, 0.1), tv, col="red", type="1")
par(oldpar)
```

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**HPLBmatrix** 

Pairwise Total Variation Distance Lower Bound Matrix for the Multi-Class Setting

#### **Description**

Pairwise Total Variation Distance Lower Bound Matrix for the Multi-Class Setting

## Usage

```
HPLBmatrix(
  labels,
  ordering.array,
  alpha = 0.05,
  computation.type = "non-optimized",
  seed = 0,
  ...
)
```

#### **Arguments**

labels a numeric vector value. The labels of the classes, should be encoded in [0,nclass1].

ordering.array a numeric array of size (nclass, nclass, nobs) such that the value (i,j,k) represents a propensity of being of class j instead of i for observation k.

alpha a numeric value. The type-I error level.

computation.type

a character value. For the moment only "non-optimized" (default) available.

seed an integer value. The seed for reproducility.

... additional parameters to be passed to the HPLB function.

#### Value

a numeric matrix of size (nclass, nclass) giving the matrix of pairwise total variation lower bounds.

#### Author(s)

Loris Michel, Jeffrey Naef

#### References

L. Michel, J. Naef and N. Meinshausen (2020). High-Probability Lower Bounds for the Total Variation Distance

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## Examples

```
# iris example
require(HPLB)
require(ranger)
# training a multi-class classifier on iris and getting tv lower bounds between classes
data("iris")
ind.train <- sample(1:nrow(iris), size = nrow(iris)/2, replace = FALSE)</pre>
rf <- ranger(Species~., data = iris[ind.train, ], probability = TRUE)</pre>
preds <- predict(rf, iris[-ind.train,])$predictions</pre>
# creating the ordering array based on prediction differences
ar <- array(dim = c(3, 3, nrow(preds)))</pre>
for (i in 1:3) {
 for (j in 1:3) {
  ar[i,j,] <- preds[,j] - preds[,i]</pre>
}
# encoding the class response
y <- factor(iris$Species)</pre>
levels(y) <- c(0,1,2)
y <- as.numeric(y)-1</pre>
# getting the lower bound matrix
tvhat.iris <- HPLBmatrix(labels = y[-ind.train], ordering.array = ar)</pre>
tvhat.iris
```

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