

# Package ‘HHG’

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**Type** Package

**Title** Heller-Heller-Gorfine Tests of Independence and Equality of Distributions

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**Depends** R (>= 3.1.0)

**Suggests** MASS, knitr

**Description** Heller-Heller-Gorfine tests are a set of powerful statistical tests of multivariate k-sample homogeneity and independence (Heller et. al., 2013, <doi:10.1093/biomet/ass070>). For the univariate case, the package also offers implementations of the 'MinP DDP' and 'MinP ADP' tests by Heller et. al. (2016), which are consistent against all continuous alternatives but are distribution-free, and are thus much faster to apply.

**License** GPL-3

**VignetteBuilder** knitr

**NeedsCompilation** yes

**RoxygenNote** 5.0.1

**Imports** Rcpp (>= 0.12.9), rmarkdown

**LinkingTo** Rcpp

**SystemRequirements** C++11

**URL** <https://github.com/barakbri/HHG>

**BugReports** <https://github.com/barakbri/HHG/issues>

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HHG-package	<i>Heller-Heller-Gorfine (HHG) Tests of Independence and Equality of Distributions</i>
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## Description

This R package implements the permutation test of independence between two random vectors of arbitrary dimensions, and equality of two or more multivariate distributions, introduced in Heller et al. (2013), as well as the distribution-free tests of independence and equality of distribution between two univariate random variables introduced in Heller et al. (2016).

## Details

Package: HHG  
 Type: Package  
 Version: 2.3.3  
 Date: 2019-05-06  
 License: GPL-2

The package contains six major functions:

[hhg.test](#) - the permutation test for independence of two multivariate (or univariate) vectors.

`hhg.test.k.sample` - the permutation test for equality of a multivariate (or univariate) distribution across K groups.

`hhg.test.2.sample` - the permutation test for equality of a multivariate (or univariate) distribution across 2 groups.

`hhg.univariate.ind.combined.test` - the distribution-free test for independence of two univariate random variables (due to the computational complexity of this function, for large sample sizes we recommend the atom based test `Fast.independence.test` instead).

`hhg.univariate.ks.combined.test` - the distribution-free test for equality of a univariate distribution across K groups.

`Fast.independence.test` - the atom based distribution-free test for independence of two univariate random variables, which is computationally efficient for large data sets (recommended for sample sizes greater than 100).

See `vignette('HHG')` for additional information.

### Author(s)

Barak Brill & Shachar Kaufman, based in part on an earlier implementation of the original HHG test by Ruth Heller <ruheller@post.tau.ac.il> and Yair Heller <heller.yair@gmail.com>. Maintainer: Barak Brill <barakbri@mail.tau.ac.il>

### References

Heller, R., Heller, Y., and Gorfine, M. (2013). A consistent multivariate test of association based on ranks of distances. *Biometrika*, 100(2), 503-510.

Heller, R., Heller, Y., Kaufman S., Brill B., & Gorfine, M. (2016). Consistent Distribution-Free K-Sample and Independence Tests for Univariate Random Variables, *JMLR* 17(29):1-54 <https://www.jmlr.org/papers/volume17/14-441/14-441.pdf>

Brill B., Heller Y., and Heller R. (2018) Nonparametric Independence Tests and k-sample Tests for Large Sample Sizes Using Package HHG, *R Journal* 10.1 <https://journal.r-project.org/archive/2018/RJ-2018-008/RJ-2018-008.pdf>

Brill B. (2016) Scalable Non-Parametric Tests of Independence (master's thesis). <http://primage.tau.ac.il/libraries/theses/exeng/free/2899741.pdf>

### Examples

```
## Not run:

# Some examples, for more see the vignette('HHG') and specific help pages

#####
#1. Univariate Independence Example
#####
#For (N<100):

N = 30
data = hhg.example.datagen(N, 'Parabola')
X = data[1,]
```

```

Y = data[2,]
plot(X,Y)

#For (N<100) , Option 1: Perform the ADP combined test
#using partitions sizes up to 4. see documentation for other parameters of the combined test
#(it is recommended to use mmax >= 4, or the default parameter for large data sets)
combined = hhg.univariate.ind.combined.test(X,Y,nr.perm = 200,mmax=4)
combined

#For (N<100) , Option 2: Perform the hhg test:

## Compute distance matrices, on which the HHG test will be based
Dx = as.matrix(dist((X), diag = TRUE, upper = TRUE))
Dy = as.matrix(dist((Y), diag = TRUE, upper = TRUE))

hhg = hhg.test(Dx, Dy, nr.perm = 1000)

hhg

#For N>100, Fast.independence.test is the recommended option:

N_Large = 1000
data_Large = hhg.example.datagen(N_Large, 'W')
X_Large = data_Large[1,]
Y_Large = data_Large[2,]
plot(X_Large,Y_Large)

NullTable_for_N_Large_MXL_tables = Fast.independence.test.nulltable(N_Large, variant = 'ADP-EQP-ML',
nr.atoms = 30,nr.perm=200)

ADP_EQP_ML_Result = Fast.independence.test(X_Large,Y_Large, NullTable_for_N_Large_MXL_tables)

ADP_EQP_ML_Result

#####
#2. Univariate K-Sample Example
#####

N0=50
N1=50
X = c(c(rnorm(N0/2,-2,0.7),rnorm(N0/2,2,0.7)),c(rnorm(N1/2,-1.5,0.5),rnorm(N1/2,1.5,0.5)))
Y = (c(rep(0,N0),rep(1,N1)))
#plot the two distributions by group index (0 or 1)
plot(Y,X)

#Option 1: Perform the distribution-free test for equality of a univariate distribution
combined.test = hhg.univariate.ks.combined.test(X,Y)
combined.test

```

```

#Option 2: Perform the permutation test for equality of distributions.

Dx = as.matrix(dist(X, diag = TRUE, upper = TRUE))

hhg = hhg.test.k.sample(Dx, Y, nr.perm = 1000)

hhg

#####
#3. Multivariate Independence Example:
#####

n=30 #number of samples
dimensions_x=5 #dimension of X matrix
dimensions_y=5 #dimension of Y matrix
X=matrix(rnorm(n*dimensions_x,mean = 0, sd = 1),nrow = n,ncol = dimensions_x) #generate noise
Y=matrix(rnorm(n*dimensions_y,mean =0, sd = 3),nrow = n,ncol = dimensions_y)

Y[,1] = Y[,1] + X[,1] + 4*(X[,1])^2 #add in the relations
Y[,2] = Y[,2] + X[,2] + 4*(X[,2])^2

#compute the distance matrix between observations.
#User may use other distance metrics.
Dx = as.matrix(dist((X)), diag = TRUE, upper = TRUE)
Dy = as.matrix(dist((Y)), diag = TRUE, upper = TRUE)

#run test
hhg = hhg.test(Dx, Dy, nr.perm = 1000)

hhg

#####
#4. Multivariate K-Sample Example
#####

#multivariate k-sample, with k=3 groups
n=100 #number of samples in each group
x1 = matrix(rnorm(2*n),ncol = 2) #group 1
x2 = matrix(rnorm(2*n),ncol = 2) #group 2
x2[,2] = 1*x2[,1] + x2[,2]
x3 = matrix(rnorm(2*n),ncol = 2) #group 3
x3[,2] = -1*x3[,1] + x3[,2]
x= rbind(x1,x2,x3)
y=c(rep(0,n),rep(1,n),rep(2,n)) #group numbers, starting from 0 to k-1

plot(x[,1],x[,2],col = y+1,xlab = 'first component of X',ylab = 'second component of X',
      main = 'Multivariate K-Sample Example with K=3 \n Groups Marked by Different Colors')

```

```
Dx = as.matrix(dist(x, diag = TRUE, upper = TRUE)) #distance matrix
hhg = hhg.test.k.sample(Dx, y, nr.perm = 1000)

hhg

## End(Not run)
```

---

Fast.independence.test

*Atoms based distribution-free test of independence*

---

### Description

Performs the atoms based distribution-free test for independence of two univariate random variables, which is computationally efficient for large data sets (recommended for sample size greater than 100).

### Usage

```
Fast.independence.test(X,Y,NullTable=NULL,mmin=2,
mmax=min(10,length(X)), variant='ADP-EQP-ML',nr.atoms = min(40,length(X)),
combining.type='MinP',score.type='LikelihoodRatio',nr.perm=200,
compress=T, compress.p0=0.001, compress.p=0.99, compress.p1=0.00001)
```

### Arguments

X	a numeric vector with observed X values.
Y	a numeric vector with observed Y values.
NullTable	The null table of the statistic, which can be downloaded from the software website or computed by the function <a href="#">Fast.independence.test.nulltable</a> .
mmin	The minimum partition size of the ranked observations, default value is 2. Ignored if NullTable is non-null.
mmax	The maximum partition size of the ranked observations, default value is the minimum between 10 and the data size.
variant	a character string specifying the partition type, must be one of "ADP-EQP" or "ADP-EQP-ML" (default). Ignored if NullTable is non-null.
nr.atoms	the number of atoms (i.e., possible split points in the data). Ignored if NullTable is non-null. The default value is the minimum between $n$ and 40.
combining.type	a character string specifying the combining type, must be one of "MinP" (default), "Fisher", or "both".
score.type	a character string specifying the score type, must be one of "LikelihoodRatio" (default), "Pearson", or "both". Ignored if NullTable is non-null.

nr.perm	The number of permutations for the null distribution. Ignored if NullTable is non-null.
compress	a logical variable indicating whether you want to compress the null tables. If TRUE, null tables are compressed: The lower compress.p part of the null statistics is kept at a compress.p0 resolution, while the upper part is kept at a compress.p1 resolution (which is finer).
compress.p0	Parameter for compression. This is the resolution for the lower compress.p part of the null distribution.
compress.p	Parameter for compression. Part of the null distribution to compress.
compress.p1	Parameter for compression. This is the resolution for the upper value of the null distribution.

## Details

This function is a smart wrapper for the [hgg.univariate.ind.combined.test](#) function, with parameters optimized for a large number of observations. The function first calls [hgg.univariate.ind.stat](#) to compute the vector of test statistics. Test statistics are the sum of log-likelihood scores, for All Derived Partitions (ADP) of the data (computed as explained in Heller et al. (2014)).

For the 'ADP-EQP-ML' variant, the base test statistics are:

$$S_{2 \times 2}, S_{2 \times 3}, S_{3 \times 2}, \dots, S_{m_{max} \times m_{max}}$$

For the 'ADP-EQP' variant, only the sum of symmetric tables (same number of cell on both axis) is considered:

$$S_{2 \times 2}, S_{3 \times 3}, S_{4 \times 4}, \dots, S_{m_{max} \times m_{max}}$$

Other variants are described in [hgg.univariate.ind.combined.test](#). The above variants are the ones to be used for a large number of observations ( $n > 100$ ).

Test functions are capable of handling large datasets by attempting a split only every  $N/nr.atoms$  observations. An atom is a sequence of observations which cannot be split when performing a partition of the data (i.e. setting `nr.atoms`, the number of sequences which cannot be split, sets the number of equidistant partition points). For the above variants, 'EQP' stands for equipartition over atoms. Brill (2016) suggests a minimum of 40 atoms, with an increase of up to 60 for alternatives which are more difficult to detect (on the expense of computational complexity. Algorithm complexity is  $O(nr.atoms^4)$ ). Very few alternatives require over 80 atoms.

The vector of  $S_{m \times l}$  statistics is then combined according to the method suggested in Heller et al. (2014). The default combining type is the minimum p-value, so the test statistic is the minimum p-value over the range of partition sizes  $m$  from `mmin` to `mmax`, where the p-value for a fixed partition size  $m$  is defined by the aggregation type and score type. The combination is done over the statistics computed by [hgg.univariate.ind.stat](#). The second type of combination method for statistics, is via a Fisher type statistic,  $-\sum \log(p_m)$  (with the sum going from `mmin` to `mmax`). The returned result may include the test statistic for the MinP combination, the Fisher combination, or both (see `comb.type`).

If the argument `NullTable` is supplied with a proper null table (constructed using

[Fast.independence.test.nulltable](#) or [hgg.univariate.ind.nulltable](#), for the data sample size), test parameters are taken from `NullTable` (`mmax`, `mmin`, `variant`, `score.type`, `nr.atoms`, ...). If `NullTable` is left NULL, a null table is generated by a call to [Fast.independence.test.nulltable](#) using the arguments supplied to this function. Null table is generated with `nr.perm` repetitions. It is

stored in the returned object, under `generated_null_table`. When testing for multiple hypotheses, one may generate only one null table (using this function or `Fast.independence.test.nulltable`), and use it many times (thus, substantially reducing computation time). Generated null tables hold the distribution of statistics for both combination types, (`comb.type=='MinP'` and `comb.type=='Fisher'`).

Null tables may be compressed, using the `compress` argument. For each of the partition sizes (i.e. `m` or `mXm`), the null distribution is held at a `compress.p0` resolution up to the `compress.p` percentile. Beyond that value, the distribution is held at a finer resolution defined by `compress.p1` (since higher values are attained when a relation exists in the data, this is required for computing the p-value accurately.)

### Value

Returns a `UnivariateStatistic` class object, with the following entries:

<code>MinP</code>	The test statistic when the combining type is "MinP".
<code>MinP.pvalue</code>	The p-value when the combining type is "MinP".
<code>MinP.m.chosen</code>	The partition size <code>m</code> for which the p-value was the smallest.
<code>Fisher</code>	The test statistic when the combining type is "Fisher".
<code>Fisher.pvalue</code>	The p-value when the combining type is "Fisher".
<code>m.stats</code>	The statistic for each <code>m</code> in the range <code>mmin</code> to <code>mmax</code> .
<code>pvalues.of.single.m</code>	The p-values for each <code>m</code> in the range <code>mmin</code> to <code>mmax</code> .
<code>generated_null_table</code>	The null table object. Null if <code>NullTable</code> is non-null.
<code>stat.type</code>	"Independence-Combined"
<code>variant</code>	a character string specifying the partition type used in the test, one of "ADP" or "DDP".
<code>aggregation.type</code>	"sum" or the aggregation type used by <code>NullTable</code>
<code>score.type</code>	a character string specifying the score type used in the test, one of "LikelihoodRatio" or "Pearson".
<code>mmax</code>	The maximum partition size of the ranked observations used for MinP or Fisher test statistic.
<code>mmin</code>	The minimum partition size of the ranked observations used for MinP or Fisher test statistic.
<code>w.sum</code>	The minimum number of observations in a partition, only relevant for <code>type="Independence"</code> , <code>aggregation.type="Sum"</code> and <code>score.type="Pearson"</code> .
<code>w.max</code>	The minimum number of observations in a partition, only relevant for <code>type="Independence"</code> , <code>aggregation.type="Max"</code> and <code>score.type="Pearson"</code> .
<code>nr.atoms</code>	The input <code>nr.atoms</code> .

### Author(s)

Barak Brill



## References

Heller, R., Heller, Y., Kaufman S., Brill B., & Gorfine, M. (2016). Consistent Distribution-Free K-Sample and Independence Tests for Univariate Random Variables, *JMLR* 17(29):1-54 <https://www.jmlr.org/papers/volume17/14-441/14-441.pdf>

Brill B., Heller Y., and Heller R. (2018) Nonparametric Independence Tests and k-sample Tests for Large Sample Sizes Using Package HHG, *R Journal* 10.1 <https://journal.r-project.org/archive/2018/RJ-2018-008/RJ-2018-008.pdf>

Brill B. (2016) Scalable Non-Parametric Tests of Independence (master's thesis) <http://primage.tau.ac.il/libraries/theses/exeng/free/2899741.pdf>

## Examples

```
## Not run:

N_Large = 1000
data_Large = hhg.example.datagen(N_Large, 'W')
X_Large = data_Large[1,]
Y_Large = data_Large[2,]
plot(X_Large,Y_Large)

NullTable_for_N_Large_MXL_tables = Fast.independence.test.nulltable(N_Large,
variant = 'ADP-EQP-ML', nr.atoms = 30,nr.perm=200)

ADP_EQP_ML_Result = Fast.independence.test(X_Large,Y_Large,
NullTable_for_N_Large_MXL_tables)

ADP_EQP_ML_Result

#the null distribution depends only on the sample size, so the same
#null table can be used for testing different hypotheses with the same sample size.
#For example, for another data set with N_Large sample size:
data_Large = hhg.example.datagen(N_Large, 'Circle')
X_Large = data_Large[1,]
Y_Large = data_Large[2,]
plot(X_Large,Y_Large)

#The MinP combining method p-values may be reported:
ADP_EQP_ML_Result = Fast.independence.test(X_Large,Y_Large,
NullTable_for_N_Large_MXL_tables,
  combining.type='MinP')
ADP_EQP_ML_Result

#or both MinP and Fisher combining methods p-values:
ADP_EQP_ML_Result = Fast.independence.test(X_Large,Y_Large,
NullTable_for_N_Large_MXL_tables,
  combining.type='Both')
ADP_EQP_ML_Result
```

```
## End(Not run)
```

---

```
Fast.independence.test.nulltable
```

*Null tables for the atoms based distribution-free test of independence.*

---

## Description

Functions for creating null table objects for the atoms based omnibus distribution-free test of independence between two univariate random variables.

## Usage

```
Fast.independence.test.nulltable(n, mmin=2, mmax=min(10, n),
  variant = 'ADP-EQP-ML', nr.atoms = min(40, n),
  score.type='LikelihoodRatio', nr.perm=200, compress=T,
  compress.p0=0.001, compress.p=0.99, compress.p1=0.000001)
```

## Arguments

<code>n</code>	The sample size
<code>mmin</code>	The minimum partition size of the ranked observations, default value is 2.
<code>mmax</code>	The maximum partition size of the ranked observations, default value is the minimum between 10 and the data size.
<code>variant</code>	a character string specifying the partition type, must be one of "ADP-EQP", "ADP-EQP-ML" (default).
<code>nr.atoms</code>	the number of atoms (i.e., possible split points in the data). Ignored if <code>NullTable</code> is non-null. The default value is the minimum between $n$ and 40.
<code>score.type</code>	a character string specifying the score type, must be one of "LikelihoodRatio" (default), "Pearson", or "both".
<code>nr.perm</code>	The number of permutations for the null distribution.
<code>compress</code>	a logical variable indicating whether you want to compress the null tables. If TRUE, null tables are compressed: The lower <code>compress.p</code> part of the null statistics is kept at a <code>compress.p0</code> resolution, while the upper part is kept at a <code>compress.p1</code> resolution (which is finer).
<code>compress.p0</code>	Parameter for compression. This is the resolution for the lower <code>compress.p</code> part of the null distribution.
<code>compress.p</code>	Parameter for compression. Part of the null distribution to compress.
<code>compress.p1</code>	Parameter for compression. This is the resolution for the upper value of the null distribution.

## Details

In order to compute the null distributions for a test statistic (with a specific aggregation and score type, and all partition sizes), the only necessary information is the sample size, since the test statistic is distribution-free. The accuracy of the quantiles of the null distribution depend on the number of replicates used for constructing the null tables. The necessary accuracy depends on the threshold used for rejection of the null hypotheses.

This function creates an object for efficiently storing the null distribution of the test statistics. Generated null tables hold the null distribution of statistics for the two combination types, i.e. for `comb.type` value ('MinP' and 'Fisher'), as well as for fixed partition sizes.

Variant types "ADP-EQP" and "ADP-EQP-ML", are the atom-based generalizations of the "ADP" and "ADP-ML". EQP type variants reduce calculation time by summing over a subset of partitions, where a split between cells may be performed only every  $n/nr.atoms$  observations. This allows for a complexity of  $O(nr.atoms^4)$ . These variants are only available for `aggregation.type=='sum'` type aggregation.

Null tables may be compressed, using the `compress` argument. For each of the partition sizes, the null distribution is held at a `compress.p0` resolution up to the `compress.p` percentile. Beyond that value, the distribution is held at a finer resolution defined by `compress.p1` (since higher values are attained when a relation exists in the data, this is required for computing the p-value accurately in the tail of the null distribution).

## Value

`m.stats` If `keep.simulation.data= TRUE`, `m.stats` a matrix with `nr.replicates` rows and `mmax-mmin+1` columns of null test statistics.

`univariate.object` A useful format of the null tables for computing p-values efficiently.

## Author(s)

Barak Brill.

## References

Heller, R., Heller, Y., Kaufman S., Brill B., & Gorfine, M. (2016). Consistent Distribution-Free K-Sample and Independence Tests for Univariate Random Variables, *JMLR* 17(29):1-54 <https://www.jmlr.org/papers/volume17/14-441/14-441.pdf>

Brill B., Heller Y., and Heller R. (2018) Nonparametric Independence Tests and k-sample Tests for Large Sample Sizes Using Package HHG, *R Journal* 10.1 <https://journal.r-project.org/archive/2018/RJ-2018-008/RJ-2018-008.pdf>

Brill B. (2016) Scalable Non-Parametric Tests of Independence (master's thesis) <http://primage.tau.ac.il/libraries/theses/exeng/free/2899741.pdf>

## Examples

```
## Not run:
```

```

N_Large = 1000
data_Large = hhg.example.datagen(N_Large, 'W')
X_Large = data_Large[1,]
Y_Large = data_Large[2,]
plot(X_Large,Y_Large)

NullTable_for_N_Large_MXM_tables = Fast.independence.test.nulltable(N_Large,
variant = 'ADP-EQP', nr.atoms = 30,nr.perm=200)
NullTable_for_N_Large_MXL_tables = Fast.independence.test.nulltable(N_Large,
variant = 'ADP-EQP-ML', nr.atoms = 30,nr.perm=200)

ADP_EQP_Result = Fast.independence.test(X_Large,Y_Large,
NullTable_for_N_Large_MXM_tables)
ADP_EQP_ML_Result = Fast.independence.test(X_Large,Y_Large,
NullTable_for_N_Large_MXL_tables)

ADP_EQP_Result
ADP_EQP_ML_Result

#null distribution depends only on data size (length(X)),
#so same null table can be used many times.
#For example, another data set:
data_Large = hhg.example.datagen(N_Large, 'Circle')
X_Large = data_Large[1,]
Y_Large = data_Large[2,]
plot(X_Large,Y_Large)

#you may use Fisher type scores:
ADP_EQP_Result = Fast.independence.test(X_Large,Y_Large,
NullTable_for_N_Large_MXM_tables, combining.type='Fisher')
#or both MinP and Fisher:
ADP_EQP_ML_Result = Fast.independence.test(X_Large,Y_Large,
NullTable_for_N_Large_MXL_tables, combining.type='Both')

ADP_EQP_Result
ADP_EQP_ML_Result

## End(Not run)

```

---

HHG

*Heller-Heller-Gorfine Tests of Independence and Equality of Distributions*


---

### Description

These functions perform Heller-Heller-Gorfine (HHG) tests. Implemented are tests of independence between two random vectors (x and y) and tests of equality of 2 or more multivariate distributions.

**Usage**

```
hhg.test(Dx, Dy, ties = T, w.sum = 0, w.max = 2, nr.perm = 10000,
         is.sequential = F, seq.total.nr.tests = 1,
         seq.alpha.hyp = NULL, seq.alpha0 = NULL, seq.beta0 = NULL, seq.eps = NULL,
         nr.threads = 0, tables.wanted = F, perm.stats.wanted = F)
```

```
hhg.test.k.sample(Dx, y, w.sum = 0, w.max = 2, nr.perm = 10000,
                 is.sequential = F, seq.total.nr.tests = 1,
                 seq.alpha.hyp = NULL, seq.alpha0 = NULL, seq.beta0 = NULL, seq.eps = NULL,
                 nr.threads = 0, tables.wanted = F, perm.stats.wanted = F)
```

```
hhg.test.2.sample(Dx, y, w.sum = 0, w.max = 2, nr.perm = 10000,
                 is.sequential = F, seq.total.nr.tests = 1,
                 seq.alpha.hyp = NULL, seq.alpha0 = NULL, seq.beta0 = NULL, seq.eps = NULL,
                 nr.threads = 0, tables.wanted = F, perm.stats.wanted = F)
```

**Arguments**

<code>Dx</code>	a symmetric matrix of doubles, where element $[i, j]$ is a norm-based distance between the $i$ 'th and $j$ 'th $x$ samples.
<code>Dy</code>	same as <code>Dx</code> , but for distances between $y$ 's (the user may choose any norm when computing <code>Dx</code> , <code>Dy</code> ).
<code>y</code>	a numeric or factor vector, whose values or levels are in $(0, 1, \dots, K - 1)$ , for performing $K$ -sample tests (including $K = 2$ ).
<code>ties</code>	a boolean specifying whether ties in <code>Dx</code> and/or <code>Dy</code> exist and are to be properly handled (requires more computation).
<code>w.sum</code>	minimum expected frequency taken into account when computing the <code>sum.chisq</code> statistic (must be non-negative, contribution of tables having cells with smaller values will be truncated to zero).
<code>w.max</code>	minimum expected frequency taken into account when computing the <code>max.chisq</code> statistic (must be non-negative, contribution of tables having cells with smaller values will be truncated to zero).
<code>nr.perm</code>	number of permutations from which a p-value is to be estimated (must be non-negative). Can be specified as zero if only the observed statistics are wanted, without p-values. The actual number of permutations used may be slightly larger when using multiple processing cores. A Wald sequential probability ratio test is optionally implemented, which may push the p-value to 1 and stop permuting if it becomes clear that it is going to be high. See Details below.
<code>is.sequential</code>	boolean flag specifying whether Wald's sequential test is desired (see Details), otherwise a simple Monte-Carlo computation of <code>nr.perm</code> permutations is performed. When this argument is TRUE, either <code>seq.total.nr.tests</code> or ( <code>seq.alpha.hyp</code> , <code>seq.alpha0</code> , <code>seq.beta0</code> , <code>seq.eps</code> ) must be supplied by the user.
<code>seq.total.nr.tests</code>	the total number of hypotheses in the family of hypotheses simultaneously tested. When this optional argument is supplied, it is used to derive default values for

the parameters of the Wald sequential test. The default derivation is done assuming a nominal 0.05 FDR level, and sets:

`seq.alpha.hyp = 0.05 / max(1, log(seq.total.nr.tests))`, `seq.alpha0 = 0.05`, `seq.beta0 = min(0.01, 0.05 / seq.total.nr.tests)`, `seq.eps = 0.01`. Alternatively, one can specify their own values for these parameters using the following arguments.

<code>seq.alpha.hyp</code>	the nominal test size for this single test within the multiple testing procedure.
<code>seq.alpha0</code>	the nominal test size for testing the side null hypothesis of $p\text{-value} > \text{seq.alpha.hyp}$ .
<code>seq.beta0</code>	one minus the power for testing the side null hypothesis of $p\text{-value} > \text{seq.alpha.hyp}$ .
<code>seq.eps</code>	approximation margin around <code>seq.alpha.hyp</code> that defines the $p\text{-value}$ regions for the side null $p > \text{seq.alpha.hyp} * (1 + \text{seq.eps})$ and side alternative $p < \text{seq.alpha.hyp} * (1 - \text{seq.eps})$ .
<code>nr.threads</code>	number of processing cores to use for $p\text{-value}$ permutation. If left as zero, will try to use all available cores.
<code>tables.wanted</code>	boolean flag determining whether to output detailed local 2x2 contingency tables.
<code>perm.stats.wanted</code>	boolean flag determining whether to output statistics values computed for all permutations (representing null distributions).

## Details

The HHG test (Heller et al., 2013) is a powerful nonparametric test for association (or, alternatively, independence) between two random vectors (say,  $x$  and  $y$ ) of arbitrary dimensions. It is consistent against virtually any form of dependence, and has been shown to offer significantly more power than alternative approaches in the face of simple, and, more importantly, complex high-dimensional dependence. The test relies on norm-based distance metrics in  $x$  and (separately) in  $y$ . The choice of metric for either variable is up to the user (e.g. Euclidean, Mahalanobis, Manhattan, or whatever is appropriate for the data). The general implementation in `hhg.test` takes the distance matrices computed on an observed sample,  $D_x$  and  $D_y$ , and starts from there.

`hhg.test.k.sample` and `hhg.test.2.sample` are optimized special cases of the general test, where  $y$  is a partition of samples in  $x$  to  $K$  or 2 groups, respectively.

When enabled by `is.sequential`, Wald's sequential test is implemented as suggested by Fay et al. (2007) in order to reduce the  $O(\text{nr.perm} * n^2 * \log(n))$  computational complexity of the permutation test to a more manageable size. Especially when faced with high multiplicity, say  $M$  simultaneous tests, the necessary number of iterations may be quite large. For example, if it is desired to control the family-wise error rate (FWER) at level  $\alpha$  using the Bonferroni correction, one needs a  $p\text{-value}$  of  $\alpha / M$  to establish significance. This seems to suggest that the minimum number of permutations required is  $\text{nr.perm} = M / \alpha$ . However, if it becomes clear after a smaller number of permutations that the null cannot be rejected, no additional permutations are needed, and the  $p\text{-value}$  can be conservatively estimated as 1. Often, only a handful of hypotheses in a family are expected to be non-null. In this case the number of permutations for testing all hypotheses using Wald's procedure is expected to be much lower than the full  $M^2 / \alpha$ .

The target significance level of the sequential test is specified in the argument `seq.alpha.hyp`. It depends on the number of hypotheses  $M$ , and the type of multiplicity correction wanted. For the Bonferroni correction, the threshold is  $\alpha / M$ . For the less conservative procedure of Benjamini

& Hochberg (1995), it is  $M1 * q / M$ , where  $q$  is the desired false discovery rate (FDR), and  $M1$  is the (unknown) number of true non-null hypotheses. Although  $M1$  is unknown, the investigator can sometimes estimate it conservatively (e.g., if at most 0.02 of the hypotheses are expected to be non-null, set  $M1 = 0.02 * M$ ).

### Value

Four statistics described in the original HHG paper are returned:

`sum.chisq` - sum of Pearson chi-squared statistics from the 2x2 contingency tables considered.

`sum.lr` - sum of likelihood ratio ("G statistic") values from the 2x2 tables.

`max.chisq` - maximum Pearson chi-squared statistic from any of the 2x2 tables.

`max.lr` - maximum G statistic from any of the 2x2 tables.

If `nr.perm > 0`, then estimated permutation p-values are returned as:

`perm.pval.hhg.sc`, `perm.pval.hhg.sl`, `perm.pval.hhg.mc`, `perm.pval.hhg.ml`

In order to give information that may help localize where in the support of the distributions of  $x$  and  $y$  there is departure from independence, if `tables.wanted` is true, the 2x2 tables themselves are provided in:

`extras.hhg.tbls`

This is a  $n^2$  by 4 matrix, whose columns are  $A_{11}$ ,  $A_{12}$ ,  $A_{21}$ ,  $A_{22}$  as denoted in the original HHG paper. Row  $r$  of the matrix corresponds to  $S_{ij}$  in the same paper, where  $i = 1 + \text{floor}((r - 1) / n)$ , and  $j = 1 + ((r - 1) \% n)$ . Since  $S_{ij}$  is never computed for  $i == j$ , rows  $(0 : (n - 1)) * n + (1 : n)$  contain NAs on purpose. The only other case where NAs will occur are for the 2 and K-sample tests, where only one table is given for any x-tied samples (the other tables at indices with the same  $x$  value are redundant).

Finally, as a means of estimating the null distributions of computed statistics, if `perm.stats.wanted` is true, the statistics computed for every permutation of the data performed during testing is outputted as:

`extras.perm.stats`

A data.frame with one variable per statistic and one sample per permutation.

`n` - The sample size

`y` - Group sizes for `hhg.test.2.sample` and `hhg.test.k.sample` tests.

`stat.type` - String holding the type of test used: `'hhg.test'`, `'hhg.test.2.sample'` or `'hhg.test.k.sample'`

### Note

The computational complexity of the test is  $n^2 * \log(n)$ , where  $n$  is the number of samples. Thus, when the sample size is large, computing the test for many permutations may take a long time.

P-values are reproducible when setting `set.seed(seed_value)` before performing the permutation test (also when computation is done in multithread). This feature is currently implemented only in `hhg.test` and not in `hhg.test.k.sample` and `hhg.test.2.sample`.

### Author(s)

Shachar Kaufman, based in part on an earlier version by Ruth Heller and Yair Heller.

## References

Heller, R., Heller, Y., & Gorfine, M. (2013). A consistent multivariate test of association based on ranks of distances. *Biometrika*, 100(2), 503-510.

Fay, M., Kim., H., & Hachey, M. (2007). On using truncated sequential probability ratio test boundaries for Monte Carlo implementation of hypothesis tests. *Journal of Computational and Graphical Statistics*, 16(4), 946-967.

Benjamini, Y., & Hochberg, Y. (1995). Controlling the false discovery rate: a practical and powerful approach to multiple testing. *Journal of the Royal Statistical Society Series B*, 57, 289-300.

## Examples

```
## Not run:
## 1. The test of independence

## 1.1. A non-null univariate example

## Generate some data from the Circle example
n = 50
X = hhg.example.datagen(n, 'Circle')
plot(X[1,], X[2,])

## Compute distance matrices, on which the HHG test will be based
Dx = as.matrix(dist((X[1,]), diag = TRUE, upper = TRUE))
Dy = as.matrix(dist((X[2,]), diag = TRUE, upper = TRUE))

## Compute HHG statistics, and p-values using 1000 random permutations
set.seed(1) #set the seed for the random permutations
hhg = hhg.test(Dx, Dy, nr.perm = 1000)

## Print the statistics and their permutation p-value

hhg

## 1.2. A null univariate example

n = 50
X = hhg.example.datagen(n, '4indclouds')

Dx = as.matrix(dist((X[1,]), diag = TRUE, upper = TRUE))
Dy = as.matrix(dist((X[2,]), diag = TRUE, upper = TRUE))

set.seed(1) #set the seed for the random permutations
hhg = hhg.test(Dx, Dy, nr.perm = 1000)

hhg

## 1.3. A multivariate example
library(MASS)

n = 50
p = 5
```



```

x = t(mvnrnorm(n, rep(0, p), diag(1, p)))
y = log(x ^ 2)
Dx = as.matrix(dist((t(x)), diag = TRUE, upper = TRUE))
Dy = as.matrix(dist((t(y)), diag = TRUE, upper = TRUE))

set.seed(1) #set the seed for the random permutations
hhg = hhg.test(Dx, Dy, nr.perm = 1000)

hhg

## 2. The k-sample test

n = 50
D = hhg.example.datagen(n, 'FourClassUniv')
Dx = as.matrix(dist(D$x, diag = TRUE, upper = TRUE))

hhg = hhg.test.k.sample(Dx, D$y, nr.perm = 1000)

hhg

## End(Not run)

```

---

hhg.example.datagen    *A set of example data generators used to demonstrate the HHG test.*

---

## Description

Six examples (Circle, Diamond, Parabola, 2Parabolas, W, 4indclouds) are taken from Newton's introduction to the discussion of the Energy Test in *The Annals of Applied Statistics* (2009). These are simple univariate dependence structures (or independence, in the latter case) used to demonstrate the tests of independence. The remaining examples (TwoClassUniv, FourClassUniv, TwoClassMultiv) generate data suitable for demonstrating the k-sample test (and in particular, the two-sample test).

It has been pointed out by Pierre Lafaye de Micheaux (private correspondence) that sampling should replace equidistant points in the data generation functions of the 2Parabolas, W, and Circle relationships (i.e., use  $x = \text{runif}(n, -1, 1)$  instead of  $x = \text{seq}(-1, 1, \text{length} = n)$ ). The power resulting from this modification is close to the original power. We did not implement this change in order to preserve the reproducibility of the results reported in paper "A consistent multivariate test of association based on ranks of distances", *Biometrika* (2016) 103 (1): 35-47.

One can see `HHG::hhg.example.datagen` and `HHG:::datagenW` (for example) for the source code for the data generation procedure.

## Usage

```
hhg.example.datagen(n, example)
```

**Arguments**

n	The desired sample size
example	The choice of example

**Value**

For `example` in {Circle, Diamond, Parabola, 2Parabolas, W, and 4indclouds}, a matrix of two rows is returned, one row per variable. Columns are i.i.d. samples. Given these data, we would like to test whether the two variables are statistically independent. Except for the 4indclouds case, all examples in fact have variables that are dependent. When `example` is one of {TwoClassUniv, FourClassUniv, TwoClassMultiv}, a list is returned with elements `x` and `y`. `y` is a vector with values either 0 or 1 (for TwoClassUniv and TwoClassMultiv) or in 0:3 for (for FourClassUniv). `x` is a real valued random variable (TwoClassUniv and FourClassUniv) or vector (TwoClassMultiv) which is not independent of `y`.

**Author(s)**

Shachar Kaufman and Ruth Heller

**References**

Newton, M.A. (2009). Introducing the discussion paper by Szekely and Rizzo. *Annals of applied statistics*, 3 (4), 1233-1235.

**Examples**

```
X = hhg.example.datagen(50, 'Diamond')
plot(X[1,], X[2,])

X = hhg.example.datagen(50, 'FourClassUniv')
plot(X)
```

---

hhg.univariate.ind.combined.test

*Distribution-free test of independence*

---

**Description**

Performs distribution-free tests for independence of two univariate random variables.

**Usage**

```
hhg.univariate.ind.combined.test(X, Y=NULL, NullTable=NULL, mmin=2,
mmax=max(floor(sqrt(length(X))/2), 2), variant='ADP', aggregation.type='sum',
score.type='LikelihoodRatio', w.sum = 0, w.max = 2, combining.type='MinP',
nr.perm=100, nr.atoms = nr_bins_equipartition(length(X)),
compress=F, compress.p0=0.001, compress.p=0.99, compress.p1=0.000001,
keep.simulation.data=T)
```

**Arguments**

X	a numeric vector with observed X values, or the test statistic as output from <a href="#">hgg.univariate.ind.stat</a> .
Y	a numeric vector with observed Y values. Leave as Null if the input to X is the test statistic.
NullTable	The null table of the statistic, which can be downloaded from the software website or computed by the function <a href="#">hgg.univariate.ind.nulltable</a> .
mmin	The minimum partition size of the ranked observations, default value is 2. Ignored if NullTable is non-null.
mmax	The maximum partition size of the ranked observations, default value is half the square root of the number of observations. For a max aggregation.type, this parameter cannot be more than 2 for the ADP variant and 4 for DDP variant. Ignored if NullTable is non-null.
variant	a character string specifying the partition type, must be one of "ADP" (default) or "DDP", "ADP-ML", "ADP-EQP", "ADP-EQP-ML". Ignored if NullTable is non-null.
aggregation.type	a character string specifying the aggregation type, must be one of "sum" (default), "max", or "both".. Ignored if NullTable is non-null.
score.type	a character string specifying the score type, must be one of "LikelihoodRatio" (default), "Pearson", or "both". Ignored if NullTable is non-null.
w.sum	The minimum number of observations in a partition, only relevant for type="Independence", aggregation.type="Sum" and score.type="Pearson", default value 0. Ignored if NullTable is non-null.
w.max	The minimum number of observations in a partition, only relevant for type="Independence", aggregation.type="Max" and score.type="Pearson", default value 2. Ignored if NullTable is non-null.
combining.type	a character string specifying the combining type, must be one of "MinP" (default), "Fisher", or "both".
nr.perm	The number of permutations for the null distribution. Ignored if NullTable is non-null.
nr.atoms	For "ADP-EQP" and "ADP-EQP-ML" type tests, sets the number of possible split points in the data. Ignored if NullTable is non-null. The default value is the minimum between $n$ and $60 + 0.5 * \sqrt{n}$ .
compress	TRUE or FALSE. If enabled, null tables are compressed: The lower compress.p part of the null statistics is kept at a compress.p0 resolution, while the upper part is kept at a compress.p1 resolution (which is finer).
compress.p0	Parameter for compression. This is the resolution for the lower compress.p part of the null distribution.
compress.p	Parameter for compression. Part of the null distribution to compress.
compress.p1	Parameter for compression. This is the resolution for the upper value of the null distribution.
keep.simulation.data	TRUE/FALSE. If TRUE, then in addition to the sorted statistics per column, the original matrix of size nr.replicates by mmax-mmin+1 is also stored.

## Details

The test statistic and p-value of the recommended independence test between two univariate random variables in Heller et al. (2014). The default combining type in the minimum p-value, so the test statistic is the minimum p-value over the range of partition sizes  $m$  from `mmin` to `mmax`, where the p-value for a fixed partition size  $m$  is defined by the aggregation type and score type. The combination is done over the statistics computed by `hhg.univariate.ind.stat`. The second type of combination method for statistics, is via a Fisher type statistic,  $-\sum \log(p_m)$  (with the sum going from `mmin` to `mmax`). The returned result may include the test statistic for the MinP combination, the Fisher combination, or both (see `comb.type`).

If the argument `NullTable` is supplied with a proper null table (constructed using `hhg.univariate.ind.nulltable`, for the data sample size), test parameters are taken from `NullTable`:

(`mmax`, `mmin`, `variant`, `aggregation.type`, `score.type`, `nr.atoms`, ...).

If `NullTable` is left `NULL`, a null table is generated by a call to `hhg.univariate.ind.nulltable` using the arguments supplied to this function. Null table is generated with `nr.perm` repetitions. It is stored in the returned object, under `generated_null_table`. When testing for multiple hypotheses, one may generate only one null table (using this function or `hhg.univariate.ind.nulltable`), and use it many times (thus, substantially reducing computation time). Generated null tables hold the distribution of statistics for both combination types, (`comb.type=='MinP'` and `comb.type=='Fisher'`).

If `X` is supplied with a statistic (`UnivariateStatistic` object, returned by `hhg.univariate.ind.stat`), `X` must have the statistics (by  $m$ ), required by either `NullTable` or the user supplied arguments `mmin` and `mmax`. If `X` has a larger `mmax` argument than the supplied null table object,  $m$  statistics which exceed the null table's `mmax` are not taken into consideration when computing the combined statistic.

Variant types "ADP-EQP" and "ADP-EQP-ML", are the computationally efficient versions of the "ADP" and "ADP-ML". EQP type variants reduce calculation time by summing over a subset of partitions, where a split between cells may be performed only every  $n/nr.atoms$  observations. This allows for a complexity of  $O(nr.atoms^4)$ . These variants are only available for `aggregation.type=='sum'` type aggregation.

Null tables may be compressed, using the `compress` argument. For each of the partition sizes (i.e.  $m$  or  $m \times m$ ), the null distribution is held at a `compress.p0` resolution up to the `compress.p0` percentile. Beyond that value, the distribution is held at a finer resolution defined by `compress.p1` (since higher values are attained when a relation exists in the data, this is required for computing the p-value accurately.)

For large data ( $n > 100$ ), it is recommended to use `Fast.independence.test`, which is an optimized version of the `hhg.univariate.ind.stat` and `hhg.univariate.ind.combined.test` tests.

## Value

Returns a `UnivariateStatistic` class object, with the following entries:

<code>MinP</code>	The test statistic when the combining type is "MinP".
<code>MinP.pvalue</code>	The p-value when the combining type is "MinP".
<code>MinP.m.chosen</code>	The partition size $m$ for which the p-value was the smallest.
<code>Fisher</code>	The test statistic when the combining type is "Fisher".
<code>Fisher.pvalue</code>	The p-value when the combining type is "Fisher".

<code>m.stats</code>	The statistic for each <code>m</code> in the range <code>mmin</code> to <code>mmax</code> .
<code>pvalues.of.single.m</code>	The p-values for each <code>m</code> in the range <code>mmin</code> to <code>mmax</code> .
<code>generated_null_table</code>	The null table object. Null if <code>NullTable</code> is non-null.
<code>stat.type</code>	"Independence-Combined"
<code>variant</code>	a character string specifying the partition type used in the test, one of "ADP" or "DDP".
<code>aggregation.type</code>	a character string specifying the aggregation type used in the , one of "sum" or "max".
<code>score.type</code>	a character string specifying the score type used in the test, one of "LikelihoodRatio" or "Pearson".
<code>mmax</code>	The maximum partition size of the ranked observations used for MinP or Fisher test statistic.
<code>mmin</code>	The minimum partition size of the ranked observations used for MinP or Fisher test statistic.
<code>w.sum</code>	The minimum number of observations in a partition, only relevant for <code>type="Independence"</code> , <code>aggregation.type="Sum"</code> and <code>score.type="Pearson"</code> .
<code>w.max</code>	The minimum number of observations in a partition, only relevant for <code>type="Independence"</code> , <code>aggregation.type="Max"</code> and <code>score.type="Pearson"</code> .
<code>nr.atoms</code>	The input <code>nr.atoms</code> .

**Author(s)**

Barak Brill and Shachar Kaufman.

**References**

Heller, R., Heller, Y., Kaufman S., Brill B, & Gorfine, M. (2016). Consistent Distribution-Free K-Sample and Independence Tests for Univariate Random Variables, *JMLR* 17(29):1-54

Brill B. (2016) Scalable Non-Parametric Tests of Independence (master's thesis)

<http://primage.tau.ac.il/libraries/theses/exeng/free/2899741.pdf>

**Examples**

```
## Not run:

N = 35
data = hhg.example.datagen(N, 'Parabola')
X = data[1,]
Y = data[2,]
plot(X,Y)

#I) Perform MinP & Fisher Tests - without existing null tables.
```

```

#Null tables are generated by the test function.
#using partitions sizes up to 5
results = hhg.univariate.ind.combined.test(X,Y,nr.perm = 100,mmax=5)
results

#The null table can then be accessed.
generated.null.table = results$generated_null_table

#II) Perform MinP & Fisher Tests - with existing null tables.

#create null table for aggregation by summation (on ADP), with partitions sizes up to 5:
ADP.null = hhg.univariate.ind.nulltable(N,mmax=5)

#create a null table, using aggregation by summation over DDP partitions,
#with partitions sizes up to 5, over Pearson scores,
#with 1000 bootstrap repetitions.
DDP.null = hhg.univariate.ind.nulltable(N,mmax = 5,variant = 'DDP',
score.type = 'Pearson', nr.replicates = 1000)

MinP.ADP.existing.null.table = hhg.univariate.ind.combined.test(X,Y, NullTable = ADP.null)
#Results
MinP.ADP.existing.null.table

#using the other null table (DDP variant, with pearson scores):
MinP.DDP.existing.null.table = hhg.univariate.ind.combined.test(X,Y, NullTable = DDP.null)

MinP.DDP.existing.null.table

# combined test can also be performed by using the test statistic.
ADP.statistic = hhg.univariate.ind.stat(X,Y,mmax=5)
MinP.using.statistic.result = hhg.univariate.ind.combined.test(ADP.statistic,
NullTable = ADP.null)
# same result as above (as MinP.ADP.result.using.existing.null.table$MinP.pvalue)
MinP.using.statistic.result$MinP.pvalue

#III) Perform MinP & Fisher Tests - using the efficient variants for large N.

N_Large = 1000
data_Large = hhg.example.datagen(N_Large, 'W')
X_Large = data_Large[1,]
Y_Large = data_Large[2,]
plot(X_Large,Y_Large)

NullTable_for_N_Large_MXM_tables = hhg.univariate.ind.nulltable(N_Large,
variant = 'ADP-EQP',nr.atoms = 30,nr.replicates=200)
NullTable_for_N_Large_MXL_tables = hhg.univariate.ind.nulltable(N_Large,
variant = 'ADP-EQP-ML',nr.atoms = 30,nr.replicates=200)

ADP_EQP_Result = hhg.univariate.ind.combined.test(X_Large,Y_Large,
NullTable_for_N_Large_MXM_tables)
ADP_EQP_ML_Result = hhg.univariate.ind.combined.test(X_Large,Y_Large,

```

```
NullTable_for_N_Large_MXL_tables)
```

```
ADP_EQP_Result
ADP_EQP_ML_Result
```

```
## End(Not run)
```

---

```
hhg.univariate.ind.nulltable
```

*Null tables for the distribution-free test of independence*

---

### Description

Functions for creating null table objects for the omnibus distribution-free test of independence between two univariate random variables.

### Usage

```
hhg.univariate.ind.nulltable(size, mmin=2, mmax = max(floor(sqrt(size)/2), 2),
  variant = 'ADP', aggregation.type = 'sum', score.type='LikelihoodRatio',
  w.sum = 0, w.max = 2, nr.replicates=1000, keep.simulation.data=F,
  nr.atoms = nr_bins_equipartition(size),
  compress=F, compress.p0=0.001, compress.p=0.99, compress.p1=0.000001)
```

### Arguments

size	The sample size
mmin	The minimum partition size of the ranked observations, default value is 2.
mmax	The maximum partition size of the ranked observations, default value is half the square root of the number of observations.
variant	a character string specifying the partition type, must be one of "ADP" (default) or "DDP", "ADP-ML", "ADP-EQP", "ADP-EQP-ML".
aggregation.type	a character string specifying the aggregation type, must be one of "sum" (default) or "max".
score.type	a character string specifying the score type, must be one of "LikelihoodRatio" (default) or "Pearson".
w.sum	The minimum number of observations in a partition, only relevant for type="Independence", aggregation.type="Sum" and score.type="Pearson", default value 0.
w.max	The minimum number of observations in a partition, only relevant for type="Independence", aggregation.type="Max" and score.type="Pearson", default value 2.
nr.replicates	The number of permutations for the null distribution.

keep.simulation.data	TRUE/FALSE. If TRUE, then in addition to the sorted statistics per column, the original matrix of size nr.replicates by mmax-mmin+1 is also stored.
nr.atoms	For "ADP-EQP" and "ADP-EQP-ML" type tests, sets the number of possible split points in the data. The default value is the minimum between $n$ and $60+0.5*\sqrt{n}$ .
compress	TRUE or FALSE. If enabled, null tables are compressed: The lower compress.p part of the null statistics is kept at a compress.p0 resolution, while the upper part is kept at a compress.p1 resolution (which is finer).
compress.p0	Parameter for compression. This is the resolution for the lower compress.p part of the null distribution.
compress.p	Parameter for compression. Part of the null distribution to compress.
compress.p1	Parameter for compression. This is the resolution for the upper value of the null distribution.

## Details

In order to compute the null distributions for a test statistic (with a specific aggregation and score type, and all partition sizes), the only necessary information is the sample size (the test statistic is "distribution free"). The accuracy of the quantiles of the null distribution depend on the number of replicates used for constructing the null tables. The necessary accuracy depends on the threshold used for rejection of the null hypotheses.

This function creates an object for efficiently storing the null distribution of the test statistics (by partition size  $m$ ). Use the returned object, together with [hhg.univariate.ind.pvalue](#) to compute the P-value for the statistics computed by [hhg.univariate.ind.stat](#)

Generated null tables also hold the distribution of statistics for combination types (comb.type=='MinP' and comb.type=='Fisher'), used by [hhg.univariate.ind.combined.test](#).

Variant types "ADP-EQP" and "ADP-EQP-ML", are the computationally efficient versions of the "ADP" and "ADP-ML". EQP type variants reduce calculation time by summing over a subset of partitions, where a split between cells may be performed only every  $n/nr.atoms$  observations. This allows for a complexity of  $O(nr.atoms^4)$ . These variants are only available for aggregation.type=='sum' type aggregation.

Null tables may be compressed, using the compress argument. For each of the partition sizes (i.e.  $m$  or  $m \times m$ ), the null distribution is held at a compress.p0 resolution up to the compress.p percentile. Beyond that value, the distribution is held at a finer resolution defined by compress.p1 (since higher values are attained when a relation exists in the data, this is required for computing the p-value accurately in the tail of the null distribution.)

For large data ( $n > 100$ ), it is recommended to use `Fast.ADP.test`, which is an optimized version of the `hhg.univariate.ind.stat` and `hhg.univariate.ind.combined` tests. Null Tables for `Fast.ADP.test` can be constructed using `Fast.ADP.nulltable`.

## Value

m.stats	If keep.simulation.data= TRUE, m.stats a matrix with nr.replicates rows and mmax-mmin+1 columns of null test statistics.
univariate.object	A useful format of the null tables for computing p-values efficiently.



**Author(s)**

Barak Brill and Shachar Kaufman.

**References**

Heller, R., Heller, Y., Kaufman S., Brill B., & Gorfine, M. (2016). Consistent Distribution-Free K-Sample and Independence Tests for Univariate Random Variables, *JMLR* 17(29):1-54 <https://www.jmlr.org/papers/volume17/14-441/14-441.pdf>

Brill B. (2016) Scalable Non-Parametric Tests of Independence (master's thesis) <http://primage.tau.ac.il/libraries/theses/exeng/free/2899741.pdf>

**Examples**

```
## Not run:
#Testing for independence, sample size = 35
N=35

#null table for aggregation by summation (on ADP):
ADP.null = hhg.univariate.ind.nulltable(N)

#create a null table, using aggregation by summation over DDP partitions,
#with partitions sizes up to 5, over Pearson scores, with 1000 bootstrap repetitions.
DDP.null = hhg.univariate.ind.nulltable(N,mmax = 5,variant = 'DDP',
score.type = 'Pearson', nr.replicates = 1000)

#Create a null table for the ADP-EQP and ADP-EQP-ML variants,
#which are tailored for independence testing, with larger n (n>100):

N_large =200
ADP.EQP.null = hhg.univariate.ind.nulltable(N_large, variant = 'ADP-EQP',nr.atoms = 40)

#Null table for the sum of log likelihood scores over all possible M X L tables:
ADP.EQP.ML.null = hhg.univariate.ind.nulltable(N_large, variant = 'ADP-EQP-ML',
nr.atoms = 30,nr.replicates=200)

## End(Not run)
```

---

hhg.univariate.ind.pvalue

*The p-value computation for the test of independence using a fixed partition size*

---

**Description**

The p-value computation for the distribution free test of independence between two univariate random variables of Heller et al. (2016) ,using a fixed partition size m.

**Usage**

```
hhg.univariate.ind.pvalue(statistic, NullTable, m=min(statistic$mmax,4),l=m)
```

**Arguments**

statistic	The value of the computed statistic by the function <code>hhg.univariate.ind.stat</code> . The statistic object includes the score type (one of "LikelihoodRatio" or "Pearson"), and the aggregation type (one of "sum" or "max").
NullTable	The null table of the statistic, which can be downloaded from the software website ( <a href="http://www.math.tau.ac.il/~ruheller/Software.html">http://www.math.tau.ac.il/~ruheller/Software.html</a> ) or computed by the function <code>hhg.univariate.ind.nulltable</code> . See <code>vignette('HHG')</code> for a method of computing null tables on multiple cores.
m	The partition size.
l	For "ADP-ML" and "ADP-EQP-ML" test variants, sets the second parameter for the partition size.

**Details**

For the test statistic, the function extracts the fraction of observations in the null table that are at least as large as the test statistic, i.e. the p-value.

For 'DDP', 'ADP' and 'ADP-EQP' variants, the partition size is described by a single parameter  $m$  (since partition size is  $m \times m$ ). For 'ADP-ML' and 'ADP-EQP-ML' variants, partition sizes of data are of sizes  $m \times l$ , allowing for assymmetric tables.

**Value**

The p-value.

**Author(s)**

Barak Brill and Shachar Kaufman.

**References**

- Heller, R., Heller, Y., Kaufman S., Brill B, & Gorfine, M. (2016). Consistent Distribution-Free K-Sample and Independence Tests for Univariate Random Variables, *JMLR* 17(29):1-54 <https://www.jmlr.org/papers/volume17/14-441/14-441.pdf>
- Brill B. (2016) Scalable Non-Parametric Tests of Independence (master's thesis) <http://primage.tau.ac.il/libraries/theses/exeng/free/2899741.pdf>

**Examples**

```
## Not run:
N = 35
data = hhg.example.datagen(N, 'Parabola')
X = data[1,]
Y = data[2,]
```

```

plot(X,Y)

#I) Computing test statistics , with default parameters:

#statistic:
hhg.univariate.ADP.Likelihood.result = hhg.univariate.ind.stat(X,Y)
hhg.univariate.ADP.Likelihood.result

#null table:
ADP.null = hhg.univariate.ind.nulltable(N)
#pvalue:
hhg.univariate.ind.pvalue(hhg.univariate.ADP.Likelihood.result, ADP.null)

#II) Computing test statistics , with summation over Data Derived Partitions (DDP),
#using Pearson scores, and partition sizes up to 5:

#statistic:
hhg.univariate.DDP.Pearson.result = hhg.univariate.ind.stat(X,Y,variant = 'DDP',
  score.type = 'Pearson', mmax = 5)
hhg.univariate.DDP.Pearson.result

#null table:
DDP.null = hhg.univariate.ind.nulltable(N,mmax = 5,variant = 'DDP',
  score.type = 'Pearson', nr.replicates = 1000)

#pvalue , for different partition size:
hhg.univariate.ind.pvalue(hhg.univariate.DDP.Pearson.result, DDP.null, m =2)
hhg.univariate.ind.pvalue(hhg.univariate.DDP.Pearson.result, DDP.null, m =5)

#III) computing P-value for the variants used for large N:

N_Large = 1000
data_Large = hhg.example.datagen(N_Large, 'W')
X_Large = data_Large[1,]
Y_Large = data_Large[2,]
plot(X_Large,Y_Large)

NullTable_ADP_EQP = hhg.univariate.ind.nulltable(N_Large, variant = 'ADP-EQP',
  nr.atoms = 30,nr.replicates=200)
NullTable_ADP_EQP_ML = hhg.univariate.ind.nulltable(N_Large,
  variant = 'ADP-EQP-ML',nr.atoms = 30,nr.replicates=200)

ADP_EQP_result = hhg.univariate.ind.stat(X_Large,Y_Large,variant = 'ADP-EQP',
  nr.atoms =30)
ADP_EQP_ML_result = hhg.univariate.ind.stat(X_Large,Y_Large,variant='ADP-EQP-ML',
  nr.atoms = 30)

#P-value for the S_(5X5) statistic, the sum over all 5X5 partitions:
hhg.univariate.ind.pvalue(ADP_EQP_result,NullTable_ADP_EQP,m=5 )

#P-value for the S_(5X3) statistic, the sum over all 5X3 partitions:

```

```
hhg.univariate.ind.pvalue(ADP_EQP_ML_result,NullTable_ADP_EQP_ML,m=5,l=3)
```

```
## End(Not run)
```

---

```
hhg.univariate.ind.stat
```

*The independence test statistics for all partition sizes*

---

### Description

These statistics are used in the omnibus distribution-free test of independence between two univariate random variables, as described in Heller et al. (2016).

### Usage

```
hhg.univariate.ind.stat(x, y, variant = 'ADP', aggregation.type='sum',
score.type='LikelihoodRatio', mmax = max(floor(sqrt(length(x))/2), 2),
mmin = 2, w.sum = 0, w.max = 2, nr.atoms = nr_bins_equipartition(length(x)))
```

### Arguments

x	a numeric vector with observed X values (tied observations are broken at random).
y	a numeric vector with observed Y values (tied observations are broken at random).
variant	a character string specifying the partition type, must be one of "ADP" (default) or "DDP", "ADP-ML", "ADP-EQP", "ADP-EQP-ML".
aggregation.type	a character string specifying the aggregation type, must be one of "sum" (default), "max", or "both".
score.type	a character string specifying the score type, must be one of "LikelihoodRatio" (default), "Pearson", or "both".
mmax	The partition size of the ranked observations. The default size is half the square root of the number of observations
mmin	The partition size of the ranked observations. The default size is half the square root of the number of observations
w.sum	The minimum number of observations in a partition, only relevant for type="Independence", aggregation.type="Sum" and score.type="Pearson", default value 0.
w.max	The minimum number of observations in a partition, only relevant for type="Independence", aggregation.type="Max" and score.type="Pearson", default value 2.
nr.atoms	For "ADP-EQP" and "ADP-EQP-ML" type tests, sets the number of possible split points in the data. The default value is the minimum between $n$ and $60+0.5*\sqrt{n}$ .

## Details

For each partition size  $m = mmin, \dots, mmax$ , the function computes the scores in each of the partitions (according to score type), and aggregates all scores according to the aggregation type (see details in Heller et al. , 2014). If the score type is one of "LikelihoodRatio" or "Pearson", and the aggregation type is one of "sum" or "max", then the computed statistic will be in `statistic`, otherwise the computed statistics will be in the appropriate subset of `sum.chisq`, `sum.lr`, `max.chisq`, and `max.lr`. Note that if the variant is "ADP", all partition sizes are computed together in  $O(N^4)$ , so the score computational complexity is  $O(N^4)$ . For "DDP" and  $mmax > 4$ , the score computational complexity is  $O(N^4) * (mmax - mmin + 1)$ .

For the 'sum' aggregation type (default), The test statistic is the sum of log likelihood (or Pearson Chi-square) scores, of all partitions of size  $m \times m$  of the data, normalized by the number of partitions and the data size (thus, being an estimator of the Mutual Information). For the 'max' aggregation type, the test statistic is the maximum log likelihood (or Pearson Chi-square) score achieved by a partition of data of size  $m$ , normalized by the data size. For variant type "ADP-ML", the statistics calculated include not only the sum over  $m \times m$  tables (symmetric tables, same number of cells on each axis), but also assymmetric tables (i.e.  $m \times l$  tables).

Variant types "ADP-EQP" and "ADP-EQP-ML", are the computationally efficient versions of the "ADP" and "ADP-ML". EQP type variants reduce calculation time by summing over a subset of partitions, where a split between cells may be performed only every  $n/nr.atoms$  observations. This allows for a complexity of  $O(nr.atoms^4)$ . These variants are only available for `aggregation.type == 'sum'` type aggregation.

For large data ( $n > 100$ ), it is recommended to use `Fast.independence.test`, which is an optimized version of the `hgg.univariate.ind.stat` and `hgg.univariate.ind.combined.test` tests.

## Value

Returns a `UnivariateStatistic` class object, with the following entries:

<code>statistic</code>	The value of the computed statistic if the score type is one of "LikelihoodRatio" or "Pearson", and the aggregation type is one of "sum" or "max". One of <code>sum.chisq</code> , <code>sum.lr</code> , <code>max.chisq</code> , and <code>max.lr</code> .
<code>sum.chisq</code>	A vector of size $mmax - mmin + 1$ , where the $m - mmin + 1$ entry is the average over all Pearson chi-squared statistics from all the $m \times m$ contingency tables considered, divided by the total number of observations.
<code>sum.lr</code>	A vector of size $mmax - mmin + 1$ , where the $m - mmin + 1$ entry is the average over all LikelihoodRatio statistics from all the $m \times m$ contingency tables considered, divided by the total number of observations.
<code>max.chisq</code>	A vector of size $mmax - mmin + 1$ , where the $m - mmin + 1$ entry is the maximum over all Pearson chi-squared statistics from all the $m \times m$ contingency tables considered.
<code>max.lr</code>	A vector of size $mmax - mmin + 1$ , where the $m - mmin + 1$ entry is the maximum over all Pearson chi-squared statistics from all the $m \times m$ contingency tables considered.
<code>type</code>	"Independence"
<code>stat.type</code>	"Independence-Stat"

size	The sample size
score.type	The input score.type.
aggregation.type	The input aggregation.type.
mmin	The input mmin.
mmax	The input mmax.
additional	A vector with the input w.sum and w.max.
nr.atoms	The input nr.atoms.

**Author(s)**

Barak Brill and Shachar Kaufman.

**References**

Heller, R., Heller, Y., Kaufman S., Brill B, & Gorfine, M. (2016). Consistent Distribution-Free K-Sample and Independence Tests for Univariate Random Variables, JMLR 17(29):1-54 <https://www.jmlr.org/papers/volume17/14-441/14-441.pdf>

Brill B. (2016) Scalable Non-Parametric Tests of Independence (master's thesis) <http://primage.tau.ac.il/libraries/theses/exeng/free/2899741.pdf>

**Examples**

```
## Not run:
N = 35
data = hhg.example.datagen(N, 'Parabola')
X = data[1,]
Y = data[2,]
plot(X,Y)

#I) Computing test statistics , with default parameters(ADP statistic):
hhg.univariate.ADP.Likelihood.result = hhg.univariate.ind.stat(X,Y)

hhg.univariate.ADP.Likelihood.result

#II) Computing test statistics , with summation over Data Derived Partitions (DDP),
#using Pearson scores, and partition sizes up to 5:

hhg.univariate.DDP.Pearson.result = hhg.univariate.ind.stat(X,Y,variant = 'DDP',
  score.type = 'Pearson', mmax = 5)
hhg.univariate.DDP.Pearson.result

#III) Computing test statistics, for all M X L tables:
hhg.univariate.ADP.ML.Likelihood.result = hhg.univariate.ind.stat(X,Y,
  variant='ADP-ML', mmax = 5)
```

```

hhg.univariate.ADP.ML.Likelihood.result

#IV) Computing test statistics, using efficient variants (for large data sets):
#Note : for independence testing with n>100, Fast.ADP.test is suggested
#rather than hhg.univariate.ind.stat.

N_Large = 1000
data_Large = hhg.example.datagen(N_Large, 'W')
X_Large = data_Large[1,]
Y_Large = data_Large[2,]
plot(X_Large,Y_Large)

hhg.univariate.ADP.EQP.Likelihood.result = hhg.univariate.ind.stat(X_Large
,Y_Large,variant='ADP-EQP', mmax = 20)

hhg.univariate.ADP.EQP.Likelihood.result

#note how only nr.atoms=76 are used - only 75 possible cell split locations are
#taken into consideration when computing the sum over all possible log likelihood scores.
#this can be changed using the nr.atoms argument:

hhg.univariate.ADP.EQP.Likelihood.result = hhg.univariate.ind.stat(X_Large,Y_Large,
variant='ADP-EQP',mmax = 20, nr.atoms =100)

hhg.univariate.ADP.EQP.Likelihood.result

#V) Computing the efficient sum over all MXL tables:

hhg.univariate.ADP.EQP.ML.Likelihood.result = hhg.univariate.ind.stat(X_Large,Y_Large,
variant='ADP-EQP-ML',mmax = 5)

hhg.univariate.ADP.EQP.ML.Likelihood.result

## End(Not run)

```

---

```
hhg.univariate.ks.combined.test
```

*Distribution-free K-sample tests*

---

## Description

Performs distribution-free tests for equality of a univariate distribution across K groups.

## Usage

```

hhg.univariate.ks.combined.test(X,Y=NULL,NullTable=NULL,mmin=2,
mmax=ifelse(is.null(Y),4,max(4,round(min(table(Y))/3))), aggregation.type='sum',
score.type='LikelihoodRatio',combining.type='MinP',nr.perm=1000,
variant='KSample-Variant', nr.atoms = nr_bins_equipartition(length(X)),
compress=F,compress.p0=0.001,compress.p=0.99,compress.p1=0.000001,keep.simulation.data=T)

```

**Arguments**

X	A numeric vector of data values (tied observations are broken at random), or the test statistic as output from <code>hhg.univariate.ks.stat</code> .
Y	for k groups, a vector of integers with values 0:(k-1) which specify the group each observation belongs to. Leave as Null if the input to X is the test statistic.
NullTable	The null table of the statistic, which can be downloaded from the software website or computed by the function <code>hhg.univariate.ks.nulltable</code> .
mmin	The minimum partition size of the ranked observations, default value is 2. Ignored if NullTable is non-null.
mmax	The maximum partition size of the ranked observations, default value is 1/3 the number of observations in the smallest group. Ignored if NullTable is non-null.
aggregation.type	a character string specifying the aggregation type, must be one of "sum" (default), or "max". Ignored if NullTable is non-null or X is the test statistic.
score.type	a character string specifying the score type, must be one of "LikelihoodRatio" (default), or "Pearson". Ignored if NullTable is non-null or X is the test statistic.
combining.type	a character string specifying the combining type, must be one of "MinP" (default), "Fisher", or "both".
nr.perm	The number of permutations for the null distribution. Ignored if NullTable is non-null.
variant	Default value is 'KSample-Variant'. Setting the variant to 'KSample-Equipartition' performs the K-sample tests over partitions of the data where splits between cells are at least $n/nr.atoms$ apart.
nr.atoms	If variant is 'KSample-Equipartition', this is the number of atoms (i.e., possible split points in the data). The default value is the minimum between n and $60 + 0.5 * \sqrt{n}$ .
compress	a logical variable indicating whether you want to compress the null tables. If TRUE, the lower compress.p part of the null statistics is kept at a compress.p0 resolution, while the upper part is kept at a compress.p1 resolution (which is finer).
compress.p0	Parameter for compression. This is the resolution for the lower compress.p part of the null distribution.
compress.p	Parameter for compression. Part of the null distribution to compress.
compress.p1	Parameter for compression. This is the resolution for the upper value of the null distribution.
keep.simulation.data	a logical variable indicating whether in addition to the sorted statistics per column, the original matrix of size nr.replicates by mmax-mmin+1 is also stored. Ignored if NullTable is non-null.



### Details

The function outputs test statistics and p-values of the combined omnibus distribution-free test of equality of distributions among K groups, as described in Heller et al. (2014). The test combines statistics from a range of partition sizes. The default combining type is the minimum p-value, so the test statistic is the minimum p-value over the range of partition sizes m from mmin to mmax, where the p-value for a fixed partition size m is defined by the aggregation type and score type. The second type of combination method for statistics, is via a Fisher type statistic,  $-\sum \log(p_m)$  (with the sum going from mmin to mmax). The returned result may include the test statistic for the MinP combination, the Fisher combination, or both (see comb.type).

If the argument NullTable is supplied with a proper null table (constructed using hhg.univariate.ks.nulltable, for the K groups sample sizes), then the following test parameters are taken from NullTable: (mmax, mmin, variant, aggregation.type, score.type, nr.atoms, ...).

If NullTable is left NULL, a null table is generated by a call to hhg.univariate.ks.nulltable using the arguments supplied to this function. The null table is generated with nr.perm repetitions. It is stored in the returned object generated\_null\_table. When testing for multiple hypotheses with the same group sample sizes, it is computationally efficient to generate only one null table (using this function or hhg.univariate.ks.nulltable), and use it for all hypotheses tested. Generated null tables hold the distribution of statistics for both combination types, (comb.type=='MinP' and comb.type=='Fisher').

If X is supplied with a statistic (UnivariateStatistic object, returned by hhg.univariate.ks.stat), X must have the statistics (by m), required by either NullTable or the user supplied arguments mmin and mmax. If X has a larger mmax argument than the supplied null table object, the statistics which exceed the null table's mmax are not taken into consideration when computing the combined statistic.

Variant type "KSample-Equipartition" is the atom based version of the K-sample test. Calculation time is reduced by aggregating over a subset of partitions, where a split between cells may be performed only every n/nr.atoms observations. Atom based tests are available when aggregation.type is set to 'sum' or 'max'.

Null tables may be compressed, using the compress argument. For each of the partition sizes, the null distribution is held at a compress.p0 resolution up to the compress.p percentile. Beyond that value, the distribution is held at a finer resolution defined by compress.p1 (since higher values are attained when a relation exists in the data, this is required for computing the p-value accurately in the tail of the null distribution.)

### Value

Returns a UnivariateStatistic class object, with the following entries:

MinP	The test statistic when the combining type is "MinP".
MinP.pvalue	The p-value when the combining type is "MinP".
MinP.m.chosen	The partition size m for which the p-value was the smallest.
Fisher	The test statistic when the combining type is "Fisher".
Fisher.pvalue	The p-value when the combining type is "Fisher".
m.stats	The statistic for each m in the range mmin to mmax.
pvalues.of.single.m	The p-values for each m in the range mmin to mmax.

generated_null_table	The null table object. Null if NullTable is non-null.
stat.type	"KSample-Combined"
aggregation.type	a character string specifying the aggregation type used in the , one of "sum" or "max".
score.type	a character string specifying the score type used in the test, one of "LikelihoodRatio" or "Pearson".
mmax	The maximum partition size of the ranked observations used for MinP or Fisher test statistic.
mmin	The minimum partition size of the ranked observations used for MinP or Fisher test statistic.
nr.atoms	The input nr.atoms.

### Author(s)

Barak Brill and Shachar Kaufman.

### References

Heller, R., Heller, Y., Kaufman S., Brill B., & Gorfine, M. (2016). Consistent Distribution-Free K-Sample and Independence Tests for Univariate Random Variables, *JMLR* 17(29):1-54 <https://www.jmlr.org/papers/volume17/14-441/14-441.pdf>

Brill B. (2016) Scalable Non-Parametric Tests of Independence (master's thesis) <http://primage.tau.ac.il/libraries/theses/exeng/free/2899741.pdf>

### Examples

```
## Not run:
#Two groups, each from a different normal mixture:
N0=30
N1=30
X = c(c(rnorm(N0/2, -2, 0.7), rnorm(N0/2, 2, 0.7)), c(rnorm(N1/2, -1.5, 0.5), rnorm(N1/2, 1.5, 0.5)))
Y = (c(rep(0, N0), rep(1, N1)))
plot(Y, X)

#I) Perform MinP & Fisher Tests - without existing null tables.
#Null tables are generated by the test function.

results = hhg.univariate.ks.combined.test(X, Y, nr.perm = 100)
results

#The null table can then be accessed.
generated.null.table = results$generated_null_table

#II) Perform MinP & Fisher Tests - with existing null tables.
```

```

#null table for aggregation by summation:
sum.nulltable = hhg.univariate.ks.nulltable(c(N0,N1), nr.replicates=1000)

MinP.Sm.existing.null.table = hhg.univariate.ks.combined.test(X,Y,
NullTable = sum.nulltable)

#Results
MinP.Sm.existing.null.table

# combined test can also be performed by using the test statistic.
Sm.statistic = hhg.univariate.ks.stat(X,Y)
MinP.using.statistic = hhg.univariate.ks.combined.test(Sm.statistic,
NullTable = sum.nulltable)
# same result as above
MinP.using.statistic$MinP.pvalue

#null table for aggregation by maximization:
max.nulltable = hhg.univariate.ks.nulltable(c(N0,N1), aggregation.type = 'max',
score.type='LikelihoodRatio', mmin = 2, mmax = 10, nr.replicates = 100)

#combined test using both "MinP" and "Fisher":
MinPFisher.Mm.result = hhg.univariate.ks.combined.test(X,Y,NullTable = max.nulltable ,
combining.type = 'Both')
MinPFisher.Mm.result

#III) Perform MinP & Fisher Tests for extremely large n

#Two groups, each from a different normal mixture, total sample size is 10^4:
X_Large = c(c(rnorm(2500,-2,0.7),rnorm(2500,2,0.7)),
c(rnorm(2500,-1.5,0.5),rnorm(2500,1.5,0.5)))
Y_Large = (c(rep(0,5000),rep(1,5000)))
plot(Y_Large,X_Large)

N0_large = 5000
N1_large = 5000

Sm.EQP.null.table = hhg.univariate.ks.nulltable(c(N0_large,N1_large), nr.replicates=200,
variant = 'KSample-Equipartition', mmax = 30)
Mm.EQP.null.table = hhg.univariate.ks.nulltable(c(N0_large,N1_large), nr.replicates=200,
aggregation.type='max', variant = 'KSample-Equipartition', mmax = 30)

MinPFisher.Sm.EQP.result = hhg.univariate.ks.combined.test(X_Large, Y_Large,
NullTable = Sm.EQP.null.table ,
combining.type = 'Both')
MinPFisher.Sm.EQP.result

MinPFisher.Mm.EQP.result = hhg.univariate.ks.combined.test(X_Large, Y_Large,
NullTable = Mm.EQP.null.table ,
combining.type = 'Both')
MinPFisher.Mm.EQP.result

```

```
## End(Not run)
```

---

```
hhg.univariate.ks.nulltable
```

*The K-sample test null tables for all partition sizes*

---

### Description

Functions for creating null table objects, for the omnibus distribution-free test of equality of distributions among  $K$  groups, as described in Heller et al. (2016). To be used for the p-value computation, see examples in [hhg.univariate.ks.pvalue](#).

### Usage

```
hhg.univariate.ks.nulltable(group.sizes,mmin=2,
mmax=max(4,round(min(group.sizes)/3)),variant = 'KSample-Variant',
aggregation.type='sum',score.type='LikelihoodRatio',
nr.replicates=1000,keep.simulation.data=F,
nr.atoms = nr_bins_equipartition(sum(group.sizes)),
compress=F,compress.p0=0.001,compress.p=0.99,compress.p1=0.000001)
```

### Arguments

<code>group.sizes</code>	the number of observations in each group.
<code>mmin</code>	The minimum partition size of the ranked observations, default value is 2.
<code>mmax</code>	The maximum partition size of the ranked observations, default value is 1/3 the number of observations in the smallest group.
<code>variant</code>	Default value is 'KSample-Variant'. Setting the variant to 'KSample-Equipartition' performs the $K$ -sample tests over partitions of the data where splits between cells are at least $n/nr.atoms$ apart.
<code>aggregation.type</code>	a character string specifying the aggregation type, must be one of "sum" (default) or "max".
<code>score.type</code>	a character string specifying the score type, must be one of "LikelihoodRatio" (default) or "Pearson".
<code>nr.replicates</code>	The number of permutations for the null distribution.
<code>keep.simulation.data</code>	TRUE/FALSE. If TRUE, then in addition to the sorted statistics per column, the original matrix of size <code>nr.replicates</code> by <code>mmax-mmin+1</code> is also stored.

<code>nr.atoms</code>	For <code>variant=='KSample-Equipartition'</code> type tests, sets the number of possible split points in the data. The default value is the minimum between $n$ and $60 + 0.5 * \sqrt{n}$ .
<code>compress</code>	TRUE or FALSE. If enabled, null tables are compressed: The lower <code>compress.p0</code> part of the null statistics is kept at a <code>compress.p0</code> resolution, while the upper part is kept at a <code>compress.p1</code> resolution (which is finer)..
<code>compress.p0</code>	Parameter for compression. This is the resolution for the lower <code>compress.p</code> part of the null distribution.
<code>compress.p</code>	Parameter for compression. Part of the null distribution to compress.
<code>compress.p1</code>	Parameter for compression. This is the resolution for the upper value of the null distribution.

## Details

In order to compute the null distributions for a test statistic (with a specific aggregation and score type, and all partition sizes), the only necessary information is the group sizes (the test statistic is "distribution free"). The accuracy of the quantiles of the null distribution depend on the number of replicates used for constructing the null tables. The necessary accuracy depends on the threshold used for rejection of the null hypotheses.

This function creates an object for efficiently storing the null distribution of the test statistics (by partition size  $m$ ). Use the returned object, together with [hhg.univariate.ks.pvalue](#) to compute the P-value for the statistics computed by [hhg.univariate.ks.stat](#)

Generated null tables also hold the distribution of statistics for combination types (`comb.type=='MinP'` and `comb.type=='Fisher'`), used by [hhg.univariate.ks.combined.test](#).

Variant type "KSample-Equipartition" is the computationally efficient version of the K-sample test. calculation time is reducing by aggregating over a subset of partitions, where a split between cells may be performed only every  $n/nr.atoms$  observations. This allows for a complexity of  $O(nr.atoms^2)$  (instead of  $O(n^2)$ ). Computationally efficient versions are available for `aggregation.type=='sum'` and `aggregation.type=='max'` variants.

Null tables may be compressed, using the `compress` argument. For each of the partition sizes (i.e.  $m$  or  $m \times m$ ), the null distribution is held at a `compress.p0` resolution up to the `compress.p` percentile. Beyond that value, the distribution is held at a finer resolution defined by `compress.p1` (since higher values are attained when a relation exists in the data, this is required for computing the p-value accurately.)

## Value

<code>m.stats</code>	If <code>keep.simulation.data=TRUE</code> , a matrix with <code>nr.replicates</code> rows and <code>mmax-mmin+1</code> columns of null test statistics.
<code>univariate.object</code>	A useful format of the null tables for computing p-values efficiently.

## Author(s)

Barak Brill and Shachar Kaufman.

## References

Heller, R., Heller, Y., Kaufman S., Brill B., & Gorfine, M. (2016). Consistent Distribution-Free K-Sample and Independence Tests for Univariate Random Variables, *JMLR* 17(29):1-54 <https://www.jmlr.org/papers/volume17/14-441/14-441.pdf>

Brill B. (2016) Scalable Non-Parametric Tests of Independence (master's thesis) <http://primage.tau.ac.il/libraries/theses/exeng/free/2899741.pdf>

## Examples

```
## Not run:
#Testing for the difference between two groups, each from a normal mixture:
N0=30
N1=30

#null table for aggregation by summation:
sum.nulltable = hhg.univariate.ks.nulltable(c(N0,N1), nr.replicates=100)
#default nr. of replicates is 1000,
#but may take several seconds. For illustration only, we use 100 replicates,
#but it is highly recommended to use at least 1000 in practice.

#null table for aggregation by maximization:
max.nulltable = hhg.univariate.ks.nulltable(c(N0,N1), aggregation.type = 'max',
  score.type='LikelihoodRatio', mmin = 3, mmax = 5, nr.replicates = 100)
#default nr. of replicates is 1000, but may take several seconds. For illustration only,
#we use 100 replicates, but it is highly recommended to use at least 1000 in practice.

#null tables for aggregation by summation and maximization, for large data variants:
#make sure to change mmax, such that mmax<= nr.atoms

N0_large = 5000
N1_large = 5000

Sm.EQP.null.table = hhg.univariate.ks.nulltable(c(N0_large,N1_large),
nr.replicates=200, variant = 'KSample-Equipartition', mmax = 30)
Mm.EQP.null.table = hhg.univariate.ks.nulltable(c(N0_large,N1_large),
nr.replicates=200, aggregation.type='max', variant = 'KSample-Equipartition', mmax = 30)

## End(Not run)
```

---

hhg.univariate.ks.pvalue

*The p-value computation for the K-sample problem using a fixed partition size*

---

## Description

The p-value computation for the K-sample test of Heller et al. (2016) using a fixed partition size  $m$ .

**Usage**

```
hgg.univariate.ks.pvalue(statistic, NullTable,m)
```

**Arguments**

statistic	The value of the computed statistic by the function <a href="#">hgg.univariate.ks.stat</a> . The statistic object includes the score type (one of "LikelihoodRatio" or "Pearson"), and the aggregation type (one of "sum" or "max").
NullTable	The null table of the statistic, which can be downloaded from the software website ( <a href="http://www.math.tau.ac.il/~ruheller/Software.html">http://www.math.tau.ac.il/~ruheller/Software.html</a> ) or computed by the function <a href="#">hgg.univariate.ind.nulltable</a> . See <code>vignette('HHG')</code> for a method of computing null tables on multiple cores.
m	The partition size.

**Details**

For the test statistic, the function extracts the fraction of observations in the null table that are at least as large as the test statistic, i.e. the p-value.

**Value**

The p-value.

**Author(s)**

Barak Brill Shachar Kaufman.

**References**

Heller, R., Heller, Y., Kaufman S., Brill B, & Gorfine, M. (2016). Consistent Distribution-Free K-Sample and Independence Tests for Univariate Random Variables, *JMLR* 17(29):1-54 <https://www.jmlr.org/papers/volume17/14-441/14-441.pdf>

Brill B. (2016) Scalable Non-Parametric Tests of Independence (master's thesis) <http://primage.tau.ac.il/libraries/theses/exeng/free/2899741.pdf>

**Examples**

```
## Not run:

#Two groups, each from a different normal mixture:
N0=30
N1=30
X = c(c(rnorm(N0/2,-2,0.7),rnorm(N0/2,2,0.7)),c(rnorm(N1/2,-1.5,0.5),rnorm(N1/2,1.5,0.5)))
Y = (c(rep(0,N0),rep(1,N1)))
plot(Y,X)

#I)p-value for fixed partition size using the sum aggregation type
hgg.univariate.Sm.Likelihood.result = hgg.univariate.ks.stat(X,Y)
```

```

hhg.univariate.Sm.Likelihood.result

sum.nulltable = hhg.univariate.ks.nulltable(c(N0,N1), nr.replicates=100)
#default nr. of replicates is 1000, but may take several seconds.
#For illustration only, we use 100 replicates, but it is highly recommended
#to use at least 1000 in practice.

#p-value for m=4 (the default):
hhg.univariate.ks.pvalue(hhg.univariate.Sm.Likelihood.result, sum.nulltable, m=4)

#p-value for m=2:
hhg.univariate.ks.pvalue(hhg.univariate.Sm.Likelihood.result, sum.nulltable, m=2)

#II) p-value for fixed partition size using the max aggregation type

hhg.univariate.Mm.likelihood.result = hhg.univariate.ks.stat(X,Y,aggregation.type = 'max')

hhg.univariate.Mm.likelihood.result

max.nulltable = hhg.univariate.ks.nulltable(c(N0,N1), aggregation.type = 'max',
  score.type='LikelihoodRatio', mmin = 3, mmax = 5, nr.replicates = 100)
#default nr. of replicates is 1000, but may take several seconds.
#For illustration only, we use 100 replicates,
#but it is highly recommended to use at least 1000 in practice.

#p-value for m=3:
hhg.univariate.ks.pvalue(hhg.univariate.Mm.likelihood.result, max.nulltable ,m = 3)

#p-value for m=5:
hhg.univariate.ks.pvalue(hhg.univariate.Mm.likelihood.result, max.nulltable,m = 5)

#III) p-value for sum and max aggregation type, using large data variants:

#Two groups, each from a different normal mixture, total sample size is 10^4:
X_Large = c(c(rnorm(2500,-2,0.7),rnorm(2500,2,0.7)),
c(rnorm(2500,-1.5,0.5),rnorm(2500,1.5,0.5)))
Y_Large = (c(rep(0,5000),rep(1,5000)))
plot(Y_Large,X_Large)

# for these variants, make sure to change mmax so that mmax<= nr.atoms

hhg.univariate.Sm.EQP.Likelihood.result = hhg.univariate.ks.stat(X_Large,Y_Large,
variant = 'KSample-Equipartition',mmax=30)
hhg.univariate.Mm.EQP.likelihood.result = hhg.univariate.ks.stat(X_Large,Y_Large,
aggregation.type = 'max',variant = 'KSample-Equipartition',mmax=30)

N0_large = 5000
N1_large = 5000

Sm.EQP.null.table = hhg.univariate.ks.nulltable(c(N0_large,N1_large), nr.replicates=200,

```



```

variant = 'KSample-Equipartition', mmax = 30)
Mm.EQP.null.table = hhg.univariate.ks.nulltable(c(N0_large,N1_large), nr.replicates=200,
aggregation.type='max', variant = 'KSample-Equipartition', mmax = 30)

hhg.univariate.ks.pvalue(hhg.univariate.Sm.EQP.Likelihood.result, Sm.EQP.null.table, m=5)
hhg.univariate.ks.pvalue(hhg.univariate.Mm.EQP.likelihood.result, Mm.EQP.null.table, m=5)

## End(Not run)

```

---

```
hhg.univariate.ks.stat
```

*The K-sample test statistics for all partition sizes*

---

### Description

These statistics are used in the omnibus distribution-free test of equality of distributions among K groups, as described in Heller et al. (2016).

### Usage

```

hhg.univariate.ks.stat(x, y, variant = 'KSample-Variant', aggregation.type='sum',
score.type='LikelihoodRatio', mmax = max(4, round(min(table(y))/3)), mmin=2,
nr.atoms= nr_bins_equipartition(length(x)))

```

### Arguments

x	a numeric vector of data values. Tied observations are broken at random.
y	for k groups, a vector of integers with values 0:(k-1) which specify the group each observation belongs to.
variant	Default value is 'KSample-Variant'. Setting the variant to 'KSample-Equipartition' performs the K-sample tests over partitions of the data where splits between cells are at least $n/nr.atoms$ apart.
aggregation.type	a character string specifying the aggregation type, must be one of "sum" (default), "max", or "both".
score.type	a character string specifying the score type, must be one of "LikelihoodRatio" (default), "Pearson", or "both".
mmax	The maximum partition size of the ranked observations, default value is 1/3 the number of observations in the smallest group.
mmin	The minimum partition size of the ranked observations, default value is 2.
nr.atoms	For variant=='KSample-Equipartition' type tests, sets the number of possible split points in the data. The default value is the minimum between $n$ and $60 + 0.5 * \sqrt{n}$ .

## Details

For each partition size  $m = mmin, \dots, mmax$ , the function computes the scores in each of the partitions (according to score type), and aggregates all scores according to the aggregation type (see details in Heller et al. , 2014). If the score type is one of "LikelihoodRatio" or "Pearson", and the aggregation type is one of "sum" or "max", then the computed statistic will be in `statistic`, otherwise the computed statistics will be in the appropriate subset of `sum.chisq`, `sum.lr`, `max.chisq`, and `max.lr`.

For the 'sum' aggregation type (default), The test statistic is the sum of log likelihood (or Pearson Chi-square) scores, of all partitions of size  $m$  of the data, normalized by the number of partitions and the data size (thus, being an estimator of the Mutual Information). For the 'max' aggregation type, the test statistic is the maximum log likelihood (or Pearson Chi-square) score achieved by a partition of data of size  $m$ .

Variant type "KSample-Equipartition" is the computationally efficient version of the K-sample test. calculation time is reducing by aggregating over a subset of partitions, where a split between cells may be performed only every  $n/nr.atoms$  observations. This allows for a complexity of  $O(nr.atoms^2)$  (instead of  $O(n^2)$ ). Computationly efficient versions are available for `aggregation.type=='sum'` and `aggregation.type=='max'` variants.

## Value

Returns a `UnivariateStatistic` class object, with the following entries:

<code>statistic</code>	The value of the computed statistic if the score type is one of "LikelihoodRatio" or "Pearson", and the aggregation type is one of "sum" or "max". One of <code>sum.chisq</code> , <code>sum.lr</code> , <code>max.chisq</code> , and <code>max.lr</code> .
<code>sum.chisq</code>	A vector of size $mmax - mmin + 1$ , where the $m - mmin + 1$ entry is the average over all Pearson chi-squared statistics from all the $K \times m$ contingency tables considered, divided by the total number of observations.
<code>sum.lr</code>	A vector of size $mmax - mmin + 1$ , where the $m - mmin + 1$ entry is the average over all LikelihoodRatio statistics from all the $K \times m$ contingency tables considered, divided by the total number of observations.
<code>max.chisq</code>	A vector of size $mmax - mmin + 1$ , where the $m - mmin + 1$ entry is the maximum over all Pearson chi-squared statistics from all the $K \times m$ contingency tables considered.
<code>max.lr</code>	A vector of size $mmax - mmin + 1$ , where the $m - mmin + 1$ entry is the maximum over all Pearson chi-squared statistics from all the $K \times m$ contingency tables considered.
<code>type</code>	"KSample".
<code>stat.type</code>	"KSample".
<code>size</code>	A vector of size K of the ordered group sample sizes.
<code>score.type</code>	The input <code>score.type</code> .
<code>aggregation.type</code>	The input <code>aggregation.type</code> .
<code>mmin</code>	The input <code>mmin</code> .
<code>mmax</code>	The input <code>mmax</code> .
<code>nr.atoms</code>	The input <code>nr.atoms</code> .

**Author(s)**

Barak Brill and Shachar Kaufman.

**References**

Heller, R., Heller, Y., Kaufman S., Brill B, & Gorfine, M. (2016). Consistent Distribution-Free K-Sample and Independence Tests for Univariate Random Variables, *JMLR* 17(29):1-54

Brill B. (2016) Scalable Non-Parametric Tests of Independence (master's thesis)

<http://primage.tau.ac.il/libraries/theses/exeng/free/2899741.pdf>

**Examples**

#Example of computing the test statistics for data from a two-sample problem:

#Two groups, each from a different normal mixture:

```
X = c(c(rnorm(25,-2,0.7),rnorm(25,2,0.7)),c(rnorm(25,-1.5,0.5),rnorm(25,1.5,0.5)))
```

```
Y = (c(rep(0,50),rep(1,50)))
```

```
plot(Y,X)
```

#I) Computing test statistics , with default parameters:

```
hhg.univariate.Sm.Likelihood.result = hhg.univariate.ks.stat(X,Y)
```

```
hhg.univariate.Sm.Likelihood.result
```

#II) Computing test statistics , with max aggregation type:

```
hhg.univariate.Mm.likelihood.result = hhg.univariate.ks.stat(X,Y,aggregation.type = 'max')
```

```
hhg.univariate.Mm.likelihood.result
```

#III) Computing statistics, which are computationally efficient for large data:

#Two groups, each from a different normal mixture, total sample size is  $10^4$ :

```
X_Large = c(c(rnorm(2500,-2,0.7),rnorm(2500,2,0.7)),
```

```
c(rnorm(2500,-1.5,0.5),rnorm(2500,1.5,0.5)))
```

```
Y_Large = (c(rep(0,5000),rep(1,5000)))
```

```
plot(Y_Large,X_Large)
```

# for these variants, make sure to change mmax so that  $mmax \leq nr.atoms$

```
hhg.univariate.Sm.EQP.Likelihood.result = hhg.univariate.ks.stat(X_Large,Y_Large,
variant = 'KSample-Equipartition',mmax=30)
```

```
hhg.univariate.Sm.EQP.Likelihood.result
```

```
hhg.univariate.Mm.EQP.likelihood.result = hhg.univariate.ks.stat(X_Large,Y_Large,
aggregation.type = 'max',variant = 'KSample-Equipartition',mmax=30)
```

```
hhg.univariate.Mm.EQP.likelihood.result
```

---

 hhg.univariate.nulltable.from.mstats

*Constructor of Distribution Free Null Table Using Existing Statistics*


---

## Description

This function converts null test statistics for different partition sizes into the null table object necessary for the computation of p-values efficiently.

## Usage

```
hhg.univariate.nulltable.from.mstats(m.stats,minm,maxm,type,variant,
size,score.type,aggregation.type, w.sum = 0, w.max = 2,
keep.simulation.data=F,nr.atoms = nr_bins_equipartition(sum(size)),
compress=F,compress.p0=0.001,compress.p=0.99,compress.p1=0.000001)
```

## Arguments

m.stats	A matrix with B rows and $\maxm - \minm + 1$ columns, where each row contains the test statistics for partition sizes m from $\minm$ to $\maxm$ for the sample permutation of the input sample.
minm	The minimum partition size of the ranked observations, default value is 2.
maxm	The maximum partition size of the ranked observations.
type	A character string specifying the test type, must be one of "KSample", "Independence"
variant	A character string specifying the partition type for the test of independence, must be one of "ADP", "DDP", "ADP-ML", "ADP-EQP", "ADP-EQP-ML" if type="Independence". If type="KSample", must be "KSample-Variant" or "KSample-Equipartition".
size	The sample size if type="Independence", and a vector of group sizes if type="KSample".
score.type	a character string specifying the score type, must be one of "LikelihoodRatio", or "Pearson".
aggregation.type	a character string specifying the aggregation type, must be one of "sum", or "max".
w.sum	The minimum number of observations in a partition, only relevant for type="Independence", aggregation.type="Sum" and score.type="Pearson", default value 0.
w.max	The minimum number of observations in a partition, only relevant for type="Independence", aggregation.type="Max" and score.type="Pearson", default value 2.
keep.simulation.data	TRUE/FALSE.
nr.atoms	For "ADP-EQP", "ADP-EQP-ML" and "KSample-Equipartition" type tests, sets the number of possible split points in the data
compress	TRUE or FALSE. If enabled, null tables are compressed: The lower compress.p part of the null statistics is kept at a compress.p0 resolution, while the upper part is kept at a compress.p1 resolution (which is finer)..

<code>compress.p0</code>	Parameter for compression. This is the resolution for the lower <code>compress.p</code> part of the null distribution.
<code>compress.p</code>	Parameter for compression. Part of the null distribution to compress.
<code>compress.p1</code>	Parameter for compression. This is the resolution for the upper value of the null distribution.

### Details

For finding multiple quantiles, the null table object is more efficient than a matrix of a matrix with  $B$  rows and  $\max_m - \min_m + 1$  columns, where each row contains the test statistics for partition sizes  $m$  from  $\min_m$  to  $\max_m$  for the sample permutation of the input sample.

Null tables may be compressed, using the `compress` argument. For each of the partition sizes (i.e.  $m$  or  $m \times m$ ), the null distribution is held at a `compress.p0` resolution up to the `compress.p` quantile. Beyond that value, the distribution is held at a finer resolution defined by `compress.p1` (since higher values are attained when a relation exists in the data, this is required for computing the p-value accurately.)

See `vignette('HHG')` for a section on how to use this function, for computing a null tables using multiple cores.

### Value

`m.stats` The input `m.stats` if `keep.simulation.data=TRUE`

`univariate.object` A useful format of the null tables for computing p-values efficiently..

### Author(s)

Barak Brill and Shachar Kaufman.

### References

Heller, R., Heller, Y., Kaufman S., Brill B, & Gorfine, M. (2016). Consistent Distribution-Free K-Sample and Independence Tests for Univariate Random Variables, *JMLR* 17(29):1-54

Brill B. (2016) Scalable Non-Parametric Tests of Independence (master's thesis)

<http://primage.tau.ac.il/libraries/theses/exeng/free/2899741.pdf>

### Examples

```
## Not run:

# 1. Downloading a lookup table from site
# download from site http://www.math.tau.ac.il/~ruheller/Software.html
#####
#using an already ready null table as object (for use in test functions)
#for example, ADP likelihood ratio statistics, for the independence problem,
#for sample size n=300
load('Object-ADP-n_300.Rdata') #=>null.table
```

```

#or using a matrix of statistics generated for the null distribution,
#to create your own table.
load('ADP-nullsim-n_300.Rdata') #=>mat
null.table = hhg.univariate.nulltable.from.mstats(m.stats = mat,minm = 2,
          maxm = 5,type = 'Independence', variant = 'ADP',size = 300,
          score.type = 'LikelihoodRatio',aggregation.type = 'sum')

# 2. generating an independence null table using multiple cores,
#and then compiling to object.
#####
library(parallel)
library(doParallel)
library(foreach)
library(doRNG)

#generate an independence null table
nr.cores = 4 #this is computer dependent
n = 30 #size of independence problem
nr.reps.per.core = 25
mmax =5
score.type = 'LikelihoodRatio'
aggregation.type = 'sum'
variant = 'ADP'

#generating null table of size 4*25

#single core worker function
generate.null.distribution.statistic =function(){
  library(HHG)
  null.table = matrix(NA,nrow=nr.reps.per.core,ncol = mmax-1)
  for(i in 1:nr.reps.per.core){
    #note that the statistic is distribution free (based on ranks),
    #so creating a null table (for the null distribution)
    #is essentially permuting over the ranks
    statistic = hhg.univariate.ind.stat(1:n,sample(1:n),
          variant = variant,
          aggregation.type = aggregation.type,
          score.type = score.type,
          mmax = mmax)$statistic

    null.table[i,]=statistic
  }
  rownames(null.table)=NULL
  return(null.table)
}

#parallelize over cores
cl = makeCluster(nr.cores)
registerDoParallel(cl)
res = foreach(core = 1:nr.cores, .combine = rbind, .packages = 'HHG',
          .export=c('variant','aggregation.type','score.type',
          'mmax','nr.reps.per.core','n'), .options.RNG=1234) %dorng%
          { generate.null.distribution.statistic() }
stopCluster(cl)

```

```

#the null table:
head(res)

#as object to be used:
null.table = hhg.univariate.nulltable.from.mstats(res,minm=2,
  maxm = mmax,type = 'Independence',
  variant = variant,size = n,score.type = score.type,
  aggregation.type = aggregation.type)

#using the null table, checking for dependence in a linear relation
x=rnorm(n)
y=x+rnorm(n)
ADP.test = hhg.univariate.ind.combined.test(x,y,null.table)
ADP.test$MinP.pvalue #pvalue

# 3. generating a k-sample null table using multiple cores
# and then compiling to object.
#####

library(parallel)
library(doParallel)
library(foreach)
library(doRNG)

#generate a k sample null table
nr.cores = 4 #this is computer dependent
n1 = 25 #size of first group
n2 = 25 #size of first group
nr.reps.per.core = 25
mmax =5
score.type = 'LikelihoodRatio'
aggregation.type = 'sum'

#generating null table of size 4*25

#single core worker function
generate.null.distribution.statistic =function(){
  library(HHG)
  null.table = matrix(NA,nrow=nr.reps.per.core,ncol = mmax-1)
  for(i in 1:nr.reps.per.core){
    #note that the statistic is distribution free (based on ranks),
    #so creating a null table (for the null distribution)
    #is essentially permuting over the ranks
    statistic = hhg.univariate.ks.stat(1:(n1+n2),sample(c(rep(0,n1),rep(1,n2))),
      aggregation.type = aggregation.type,
      score.type = score.type,
      mmax = mmax)$statistic

    null.table[i,]=statistic
  }
  rownames(null.table)=NULL
  return(null.table)
}

```

```

}

#parallelize over cores
cl = makeCluster(nr.cores)
registerDoParallel(cl)
res = foreach(core = 1:nr.cores, .combine = rbind, .packages = 'HHG',
              .export=c('n1','n2','aggregation.type','score.type','mmax',
                        'nr.reps.per.core'), .options.RNG=1234) %dorng%
              {generate.null.distribution.statistic()}
stopCluster(cl)

#the null table:
head(res)

#as object to be used:
null.table = hhg.univariate.nulltable.from.mstats(res,minm=2,
          maxm = mmax,type = 'KSample',
          variant = 'KSample-Variant',size = c(n1,n2),score.type = score.type,
          aggregation.type = aggregation.type)

#using the null table, checking for dependence in a case of two distinct samples
x=1:(n1+n2)
y=c(rep(0,n1),rep(1,n2))
Sm.test = hhg.univariate.ks.combined.test(x,y,null.table)
Sm.test$MinP.pvalue #pvalue

## End(Not run)

```

---

nr\_bins\_equipartition *Function for selecting the default number of atoms for equipartition*

---

### Description

Function selects the default number of atoms by sample size, being  $\min(n, 60 + 0.5 \cdot \text{floor}(\sqrt{n}))$

### Usage

```
nr_bins_equipartition(n)
```

### Arguments

n	Sample size
---	-------------

### Details

Function returns the default number of atoms (bins of sample space partition) for tests performed over a large number of observations. The default number of atoms is  $\min(n, 60 + 0.5 \cdot \text{floor}(\sqrt{n}))$ .



**Value**

Default number of atoms by sample size.

**Author(s)**

Barak Brill

**Examples**

```
nr_bins_equipartition(100)
```

---

`print.HHG.Test.Result` *Print function for result of HHG tests*

---

**Description**

Print description of for result object of HHG tests

**Usage**

```
## S3 method for class 'HHG.Test.Result'  
print(x, ...)
```

**Arguments**

<code>x</code>	result of <code>hhg.test</code> , <code>hhg.test.2.sample</code> or <code>hhg.test.k.sample</code>
<code>...</code>	Additional arguments can be sent to function. Currently not supported.

**Details**

Function prints description of results for the `hhg.test`, `hhg.test.2.sample` and `hhg.test.k.sample` functions. Displays: test statistics, pvalues (if permutations were performed) and description of sample size (also displays group sizes and equality of distribution tests).

**Value**

Does not return value. Only prints description of test statistic and results.

**Author(s)**

Barak Brill

**Examples**

```

#output for independence test
n = 50
X = hhg.example.datagen(n, '4indclouds')

Dx = as.matrix(dist((X[1,]), diag = TRUE, upper = TRUE))
Dy = as.matrix(dist((X[2,]), diag = TRUE, upper = TRUE))

hhg = hhg.test(Dx, Dy, nr.perm = 200)

#output for k-sample test
n = 50
D = hhg.example.datagen(n, 'FourClassUniv')
Dx = as.matrix(dist(D$x, diag = TRUE, upper = TRUE))

hhg = hhg.test.k.sample(Dx, D$y, nr.perm = 200)

```

---

```
print.UnivariateObject
```

*Print function for Univariate Null Table Object*

---

**Description**

Print description of univariate object.

**Usage**

```
## S3 method for class 'UnivariateObject'
print(x, ...)
```

**Arguments**

`x` A univariate null table object, of type 'UnivariateObject'  
`...` Additional arguments can be sent to function. Currently not supported.

**Details**

Function prints description of a null table object, including sample size (for the independence `hhg.univariate` statistics) or group sizes (for `hhg.univariate` statistics). Also prints statistic type in terms of `variant`, `aggregation.type` and `score.type`.

**Value**

Does not return value. Only prints description of null table object.

**Author(s)**

Barak Brill

### Examples

```
#univariate objects are found inside null tables:
nt = hhg.univariate.ks.nulltable(group.sizes = c(20,20), nr.replicates = 200)

print(nt$univariate.object)
```

---

```
print.UnivariateStatistic
      Print function for Univariate Statistic Test Object
```

---

### Description

Print description of univariate statistic result & test object.

### Usage

```
## S3 method for class 'UnivariateStatistic'
print(x, ...)
```

### Arguments

x	result of <code>hhg.univariate.ind.stat</code> , <code>hhg.univariate.ks.stat</code> , <code>hhg.univariate.ind.combined.test</code> or <code>hhg.univariate.ks.combined.test</code>
...	Additional arguments can be sent to function. Currently not supported.

### Details

Function prints description of univariate test statistic and test results. Displays: test statistics, pvalues (for the combined test function), partition sizes and type of test statistic used in terms of `variant`, `aggregation.type` and `score.type`.

### Value

Does not return value. Only prints description of test statistic and results.

### Author(s)

Barak Brill

### Examples

```
#generate statistics and test results, and print them
ind.stat = hhg.univariate.ind.stat(1:20,1:20,variant = 'ADP',
  aggregation.type = 'sum',score.type = 'both',mmax = 5)
print(ind.stat)

ks.stat = hhg.univariate.ks.stat(1:50,sample(c(rep(0,25),rep(1,25))),
```

```
    aggregation.type = 'both',score.type = 'both',mmax = 10)
print(ks.stat)

## Not run:

ind.combined = hhg.univariate.ind.combined.test(1:20,1:20,
    combining.type = 'Both',mmax = 5,nr.perm = 100)
print(ind.combined)

## End(Not run)

ks.combined = hhg.univariate.ks.combined.test(1:50,
    sample(c(rep(0,25),rep(1,25))),combining.type = 'Both')
print(ks.combined)
```

---

Yeast\_hughes

*Yeast gene expression data*

---

## Description

The data is a small subset of the full dataset from Hughes et al. (2000).

## Usage

```
data(Yeast_hughes)
```

## Format

A data frame for 10 *S. cerevisiae* (which is a common and well-studied species of Yeast) genes (stored in rows) and 300 expression observations (stored in columns). The 10 genes are those positioned 10, 20, 30, 40, 50, 60, 70, 80, 90, and 100 in chromosome I in the original data (see link below).

## Source

Functional Discovery via a Compendium of Expression Profiles. Hughes et al., Cell, 2000.

The full data is publicly available at <http://noble.gs.washington.edu/proj/microarray/Rosetta/>.

## References

Hughes et al.(2000). Functional Discovery via a Compendium of Expression Profiles. *Cell*, 102(Issue 1), P103-P126.

## Examples

```
data(Yeast_hughes)
```

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