# Package 'multivariance’ 

October 13, 2022
Title Measuring Multivariate Dependence Using Distance Multivariance
Version 2.4.1
Date 2021-10-06
Description Distance multivariance is a measure of dependence which can be used to detect and quantify dependence of arbitrarily many random vectors. The necessary functions are implemented in this packages and examples are given. It includes: distance multivariance, distance multicorrelation, dependence structure detection, tests of independence and copula versions of distance multivariance based on the Monte Carlo empirical transform. Detailed references are given in the package description, as starting point for the theoretic background we refer to:
B. Böttcher, Dependence and Dependence Structures: Estimation and Visualization Using the Unifying Concept of Distance Multivariance. Open Statistics, Vol. 1, No. 1 (2020), [doi:10.1515/stat-2020-0001](doi:10.1515/stat-2020-0001).

Depends R (>= 3.3.0)
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## Encoding UTF-8

LazyData true
Imports igraph, graphics, stats, Rcpp, microbenchmark
RoxygenNote 7.1.2
Suggests testhat
LinkingTo Rcpp
NeedsCompilation yes
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Repository CRAN
Date/Publication 2021-10-06 15:50:05 UTC

## $R$ topics documented:

multivariance-package ..... 3
anscombe.extended ..... 5
cdm ..... 6
cdms ..... 7
clean.graph ..... 8
coins ..... 9
copula.multicorrelation ..... 10
copula.multicorrelation.test ..... 11
copula.multivariance ..... 11
dependence.structure ..... 12
dep_struct_iterated_13_100 ..... 15
dep_struct_ring_15_100 ..... 16
dep_struct_several_26_100 ..... 17
dep_struct_star_9_100 ..... 17
emp.transf ..... 18
fastdist ..... 18
fastEuclideanCdm ..... 19
find.cluster ..... 19
independence.test ..... 21
layout_on_circles ..... 22
m.multivariance ..... 23
multicorrelation ..... 25
multivariance ..... 27
multivariance.pvalue ..... 29
multivariance.test ..... 29
multivariance.timing ..... 31
multivariances.all ..... 32
pearson.pvalue ..... 33
pearson.qf ..... 34
rejection.level ..... 35
resample.multivariance ..... 36
resample.pvalue ..... 37
resample.rejection.level ..... 38
sample.cdms ..... 38
sample.cols ..... 39
tetrahedron ..... 40
total.multivariance ..... 40
Index ..... 43

## Description

The multivariance package provides basic functions to calculate distance multivariance and related quantities. To test independence use multivariance.test, it provides an interface (via its arguments) to all the tests based on distance (m-/total-)multivariance. The package offers also several other functions related to distance multivariance, e.g. a detection and visualization of dependence structures dependence.structure. See below for details on the full content of the package.

## Details

Distance multivariance is a multivariate dependence measure, which can be used to detect dependencies between an arbitrary number of random vectors each of which can have a distinct dimension. The necessary functions are implemented in this package, and examples are given. For the theoretic background we refer to the papers [1,2,3,4,5,6]. Paper [3] includes a summary of the first two. It is the recommended starting point for users with an applied interest. Paper [4] is concerned with new (faster) p-value estimates for the independence tests, [5] introduces the copula versions of distance multivariance, [6] discusses the quantification of dependence using distance multicorrelations.

The (current) code is speed improved in comparison to the former releases. Certainly there is still room for improvement and development. Questions, comments and remarks are welcome: [bjoern.boettcher@tu-dresden.de](mailto:bjoern.boettcher@tu-dresden.de)
For infos on the latest changes and/or updates to the package use news (package="multivariance").
To cite this package use the standard citation for R packages, i.e., the output of citation("multivariance").

## Multivariance

multivariance computes the distance multivariance
total.multivariance computes the total distance multivariance
m.multivariance computes the m-multivariance (introduced in [3])

It might be convenient to compute these simultaneously using multivariances.all.
copula.multivariance computes the copula versions of the above (introduced in [5])
multicorrelation computes the multicorrelations (discussed specifically in [6])

## Functions to use and interpret multivariance

rejection. level computes a (conservative) rejection level for a given significance level. This can be used for a conservative interpretation of distance multivariance. The counterpart is multivariance.pvalue, which computes a conservative p-value for a given distance multivariance. Both methods are distribution-free.
resample.rejection.level and resample.pvalue are the distribution dependent versions of the above. They are approximately sharp, but computational more expensive. Any resampling is done by resample.multivariance.
Using the methods developed in [4] approximate p-value estimates are provided by pearson. pvalue. This method is much faster than the resampling method.
multivariance. test provides the corresponding tests of independence. The former provides output as common for tests in R.
cdm and cdms compute the doubly centered distance matrix and matrices, respectively. These can be used to speed up repeated computations of distance multivariance.
In [4] various methods to estimate the moments of the test statistic under H 0 were developed, these are (implicitly) implemented in this package only for the moments used in pearson. pvalue. Further and explicit functions can be added upon request. Please feel free to contact the author.
emp. transf computes the Monte Carlo empirical transform of the data. This data yields the copula version of distance multivariance. Hereto note, that values become randomized due to the "Monte Carlo empirical transform", i.e., the copula versions yield in a finite sample setting not identical values for repeated runs.
For planing of large projects or studies it might be convenient to estimate the computation time of multivariance via multivariance.timing.

## Dependence structures

dependence.structure performs the dependence structure detection algorithm as described in [3]. find.cluster is the basic building block of dependence. structure. It is recommended to use dependence. structure.

## Examples

coins and tetrahedron generate samples of pairwise independent random variables, with dependence of higher order.
dep_struct_iterated_13_100, dep_struct_ring_15_100, dep_struct_several_26_100 and dep_struct_star_9_100 are example data sets for the dependence structure detection. These might also serve as benchmark examples.
anscombe. extended provides an extension of Anscombe's Quartett. It illustrates that a large value of Pearson's correlation can occur for very different dependencies and that this is not a small-sample problem. These dependencies are at least partly differentiated by values of distance multicorrelation.

## References

[1] B. Böttcher, M. Keller-Ressel, R.L. Schilling, Detecting independence of random vectors: generalized distance covariance and Gaussian covariance. Modern Stochastics: Theory and Applications, Vol. 5, No. 3(2018) 353-383. https://www.vmsta.org/journal/VMSTA/article/127/ info
[2] B. Böttcher, M. Keller-Ressel, R.L. Schilling, Distance multivariance: New dependence measures for random vectors. The Annals of Statistics, Vol. 47, No. 5 (2019) 2757-2789. doi: 10.1214/ 18AOS1764
[3] B. Böttcher, Dependence and Dependence Structures: Estimation and Visualization using the Unifying Concept of Distance Multivariance. Open Statistics, Vol. 1, No. 1 (2020) 1-46. doi: 10.1515/ stat20200001
[4] G. Berschneider, B. Böttcher, On complex Gaussian random fields, Gaussian quadratic forms and sample distance multivariance. Preprint. https://arxiv.org/abs/1808.07280
[5] B. Böttcher, Copula versions of distance multivariance and dHSIC via the distributional transform - a general approach to construct invariant dependence measures. Statistics, (2020) 1-18. doi: 10.1080/02331888.2020.1748029
[6] B. Böttcher, Notes on the interpretation of dependence measures - Pearson's correlation, distance correlation, distance multicorrelations and their copula versions. Preprint. https://arxiv. org/abs/2004.07649
anscombe.extended Extended Anscombe's Quartett

## Description

The dataset extends 'anscombe' provided in the standard R-package 'datasets'. All examples feature the same correlation of 0.82 , but different types of dependencies. The main aim was to extend the classical examples, which have sample size 11, to larger sample sizes. This illustrates that the implied problems of Pearson's correlation are not small sample problems! Distance multicorrelation (which coincides in this case with distance correlation) yields different values for the datasets.

## Usage

anscombe.extended

## Format

list with elements:

- anscombe.extended $\$ \mathrm{~N} 11$ matrix with 11 samples for 5 examples the first 4 are the classical Anscombe Quartett, the fifth is a monoton relation which also features the same correlation.
- anscombe.extended $\$ \mathrm{~N} 100$ same as above but 100 samples
- anscombe.extended\$N1000 same as above but 1000 samples


## Details

Note: Anscombe's quartett features further identical parameters besides Pearson's correlation. The extended set is only concerned with correlation.

## References

This example was introduced in the reference [6] given on the main help page of this package: multivariance-package.

## Examples

```
# Code which generates plots of all included data:
op = par(mfrow = c(3,5),mar = c(0.5,0.5,3,0.5))
for (name in c("N11","N100","N1000")) {
    for (i in 1:5) {
        x = anscombe.extended[[name]][,2*i-1]
        y = anscombe.extended[[name]][,2*i]
        plot(x,y,main = paste0("cor = ", round(cor(x,y),2),
    "\n Mcor = ",round(multicorrelation(cbind(x,y),type = "pairwise",squared = FALSE),2),
    "\n CMcor = ",round(copula.multicorrelation(cbind(x,y),type = "pairwise", squared = FALSE), 2)),
            axes = FALSE,xlab ="",ylab = "", cex.main=1)
        # for two variables 'pairwise' coincides with
        # both values of 'total.upper.lower'.
        box()
    }
}
par(op)
```

cdm
computes a doubly centered distance matrix

## Description

computes the doubly centered distance matrix

## Usage

```
    cdm(
        x,
        normalize = TRUE,
        psi = NULL,
        p = NULL,
        isotropic = FALSE,
        external.dm.fun = NULL
    )
```


## Arguments

x
normalize
psi if it is NULL, the euclidean distance will be used. In the case of isotropic= TRUE: a real valued negative definite function of one variable (accepting vectors as arguments; returning a vector of the same length). In the case of isotropic $=$ FALSE: a real valued function of two variables (or vectors) to compute the distance of two samples based on a continuous negative definite function.
p
numeric, if it is a value between 1 and 2 then the Minkowski distance with parameter p is used.
isotropic logical, indicates if psi of the Euclidean distance matrix should be computed, i.e., if an isotropic distance should be used.
external.dm.fun
here one can supply an external function, which computes the distance matrix given $x$.

## Details

The doubly centered distance matrices are required for the computation of (total/m-) multivariance.
If normalize $=$ TRUE then the value of multivariance is comparable and meaningful. It can be compared to the rejection. level or its p-value multivariance. pvalue can be computed.
More details: If normalize = TRUE the matrix is scaled such that the multivariance based on it, times the sample size, has in the limit - in the case of independence - the distribution of an $\mathrm{L}^{\wedge} 2$ norm of a Gaussian process with known expectation.

As default the Euclidean distance is used. The parameters psi, p, isotropic and external.dm. fun can be used to select a different distance. In particular, external.dm. fun can be used to provide any function which calculates a distance matrix for the rows of a given matrix.

## References

For the theoretic background see the references given on the main help page of this package: multivariance-package.

## Examples

```
x = coins(100)
cdm(x) # fast euclidean distances
cdm(x,psi = function(x,y) sqrt(sum((x-y)^2))) # this is identical to the previous (but slower)
# the function cdm does the following three lines in a faster way
N = nrow(x)
C = diag(N) - matrix(1/N,nrow = N,ncol = N)
A = - C %*% as.matrix(stats::dist(x,method="euclidean")) %*% C #'
all(abs(A- cdm(x,normalize = FALSE)) < 10^(-12))
```

cdms
computes the doubly centered distance matrices

## Description

computes the doubly centered distance matrices

## Usage

cdms(x, vec $=1: n c o l(x)$, membership $=$ NULL, ...)

## Arguments

| $x$ | matrix, each row is a sample |
| :--- | :--- |
| vec | vector which indicates which columns are treated as one sample |
| membership | depreciated. Now use vec. |
| $\ldots$ | these are passed to cdm |

## Value

It returns a list of distance matrices.

```
clean.graph cleanup dependence structure graph
```


## Description

Given a dependence structure graph: vertices representing the multivariances of only two vertices can be turned into an edge labeled with the label of the vertex. Moreover, only subsets of the graph can be selected.

```
Usage
    clean.graph(
        g,
        only.level = NULL,
        simplify.pairs = TRUE,
        drop.label.pairs = FALSE
    )
```


## Arguments

g
only.level integer vector, if provided all edges and dependency nodes corresponding to dependence orders not given in 'only.level' are removed
simplify.pairs boolean, if true dependency nodes which are only connected to two variables are turned into edges
drop.label.pairs
boolean, if true the labels for edges indicating pairwise dependence are removed

## Details

Note: The option 'only.level' works only properly for a full dependence structure graph, in the case of a clustered dependence structure graph dependency nodes representing a cluster might be removed.

## Value

graph

## Examples

```
N = 200
y = coins(N,2)
x = cbind(y,y,y)
ds = dependence.structure(x,structure.type = "clustered")
plot(clean.graph(ds$graph))
plot(clean.graph(ds$graph,only.level = 2))
plot(clean.graph(ds$graph,only.level = 3)) # of limited use for a clustered graph,
# i.e., here the three-dependence node without edges indicates that
# all edges were connected to clusters
ds = dependence.structure(x,structure.type = "full")
plot(clean.graph(ds$graph))
plot(clean.graph(ds$graph,drop.label.pairs = TRUE))
plot(clean.graph(ds$graph,only.level = 2))
plot(clean.graph(ds$graph,only.level = 2,drop.label.pairs = TRUE))
plot(clean.graph(ds$graph,only.level = 3))
```

coins
dependence example: $k$-independent coin sampling

## Description

This function creates samples which are dependent but k-independent.

## Usage

coins(N = 1000, k = 2, type = "even")

## Arguments

N
number of samples
$k \quad$ each $k$-tuple will be independent
type one of "even" or "odd"

## Details

Throw $k$ independent fair coins. Now consider the $k+1$ events: The first shows head, the second shows head,... the $k$-th shows head, there is an even (or odd as selected via type) number of heads. Each row contains the state of these $\mathrm{k}+1$ events.

## Value

It returns the samples as rows of an $N$ by $k+1$ matrix. The columns are dependent but $k$-independent.

## References

For the theoretic background see the reference [3] given on the main help page of this package: multivariance-package.

## Examples

```
coins(200,4)
```

```
copula.multicorrelation
```

coupla versions of distance multicorrelation

## Description

Formally it is nothing but distance multicorrelation applied to the Monte Carlo emprical transform of the data. Hence its values vary for repeated runs.

## Usage

copula.multicorrelation(x, vec $=1: n \operatorname{col}(x), \ldots)$
$\operatorname{CMcor}(x$, vec $=1: n \operatorname{col}(x), \ldots)$

## Arguments

$$
\begin{array}{ll}
\mathrm{x} & \text { either a data matrix or a list of doubly centered distance matrices } \\
\text { vec } & \begin{array}{l}
\text { if } x \text { is a matrix, then this indicates which columns are treated together as one } \\
\text { sample; if } x \text { is a list, these are the indexes for which the multivariance is calcu- } \\
\text { lated. The default is all columns and all indexes, respectively. }
\end{array} \\
\ldots & \text { are passed to multicorrelation }
\end{array}
$$

## References

For the theoretic background see the reference [5] given on the main help page of this package: multivariance-package.

## See Also

multicorrelation
copula.multicorrelation.test
independence tests using the copula versions of distance multivariance

## Description

Formally it is nothing but tests for distance multivariance applied to the Monte Carlo emprical transform of the data. Hence its values vary for repeated runs.

## Usage

copula.multicorrelation.test(x, vec = 1:ncol(x), ...)

## Arguments

$x \quad$ matrix, each row is a sample
vec vector which indicates which columns are treated as one sample
... these are passed to cdm

## References

For the theoretic background see the reference [5] given on the main help page of this package: multivariance-package.
copula.multivariance copula version of distance multivariance

## Description

Formally it is nothing but distance multivariance applied to the Monte Carlo emprical transform of the data. Hence its values vary for repeated runs.

## Usage

copula.multivariance(x, vec = 1:ncol(x), type = "total", ...)

## Arguments

$x \quad$ either a data matrix or a list of doubly centered distance matrices
vec if $x$ is a matrix, then this indicates which columns are treated together as one sample; if x is a list, these are the indexes for which the multivariance is calculated. The default is all columns and all indexes, respectively.
type default: "total.lower.upper", for details and other options see below
... these are passed to cdms (which is only invoked if x is a matrix)

## References

For the theoretic background see the reference [5] given on the main help page of this package: multivariance-package.

```
dependence.structure determines the dependence structure
```


## Description

Determines the dependence structure as described in [3].

## Usage

dependence.structure(
x ,
vec = 1:ncol(x),
verbose = TRUE,
detection.aim = NULL,
type = "conservative",
structure.type = "clustered",
c.factor = 2,
list.cdm = NULL,
alpha $=0.05$,
p.adjust.method = "holm",
stop.too.many $=$ NULL,
)

## Arguments

X
vec vector, it indicates which columns are initially treated together as one sample
verbose boolean, if TRUE details are printed during the detection and whenever a cluster is newly detected the (so far) detected dependence structure is plotted.
detection.aim =NULL or a list of vectors which indicate the expected detection, see below for more details
type the method used for the detection, one of 'conservative','resample','pearson_approx' or 'consistent'
structure.type either the 'clustered' or the 'full' structure is detected
c. factor numeric, larger than 0 , a constant factor used in the case of 'type = "consistent",
list.cdm not required, the list of doubly centered distance matrices corresponding to $x$ speeds up the computation if given
alpha numeric between 0 and 1, the significance level used for the tests

```
p.adjust.method
    a string indicating the p-value adjustment for multiple testing, see p.adjust.methods
stop.too.many numeric, upper limit for the number of tested tuples. A warning is issued if it is
    used. Use stop. too.many = NULL for no limit.
... these are passed to find.cluster
```


## Details

Performs the detection of the dependence structure as described in [3]. In the clustered structure variables are clustered and treated as one variable as soon as a dependence is detected, the full structure treats always each variable separately. The detection is either based on tests with significance level alpha or a consistent estimator is used. The latter yields (in the limit for increasing sample size) under very mild conditions always the correct dependence structure (but the convergence might be very slow).
If fixed.rejection. level is not provided, the significance level alpha is used to determine which multivariances are significant using the distribution-free rejection level. As default the Holm method is used for p -value correction corresponding to multiple testing.
The resulting graph can be simplified (pairwise dependence can be represented by edges instead of vertices) using clean. graph.
Advanced: The argument detection. aim is currently only implemented for structure.type = clustered. It can be used to check, if an expected dependence structure was detected. This might be useful for simulation studies to determine the empirical power of the detection algorithm. Hereto detection. aim is set to a list of vectors which indicate the expected detected dependence structures (one for each run of find.cluster). The vector has as first element the k for which k-tuples are detected (for this aim the detection stops without success if no k-tuple is found), and the other elements, indicate to which clusters all present vertices belong after the detection, e.g. $c(3,2,2,1,2,1,1,2,1)$ expects that 3 -tuples are detected and in the graph are 8 vertices (including those representing the detected 3 dependencies), the order of the 2 's and 1 's indicate which vertices belong to which cluster. If detection. aim is provided, the vector representing the actual detection is printed, thus one can use the output with copy-paste to fix successively the expected detection aims.
Note that a failed detection might invoke the warning:

```
run$mem == detection.aim[[k]][-1] :
longer object length is not a multiple of shorter object length
```


## Value

returns a list with elements:
multivariances calculated multivariances, cdms calculated doubly centered distance matrices, graph graph representing the dependence structure, detected boolean, this is only included if a detection.aim is given, number. of. dep. tuples vector, with the number of dependent tuples for each tested order. For the full dependence structure a value of -1 indicates that all tuples of this order are already lower order dependent, a value of -2 indicates that there were more than stop. too.many tuples,
structure.type either clustered or full,
type the type of p-value estimation or consistent estimation used,
total. number. of.tests numeric vector, with the number of tests for each group of tests,
typeI. error. prob estimated probability of a type I error,
alpha significance level used if a p-value estimation procedure is used,
c. factor factor used if a consistent estimation procedure is used,
parameter. range significance levels (or 'c.factor' values) which yield the same detection result.

## References

For the theoretic background see the reference [3] given on the main help page of this package: multivariance-package.

## Examples

```
# structures for the datasets included in the package
dependence.structure(dep_struct_several_26_100)
dependence.structure(dep_struct_star_9_100)
dependence.structure(dep_struct_iterated_13_100)
dependence.structure(dep_struct_ring_15_100)
# basic examples:
x = coins(100) # 3-dependent
dependence.structure(x)
colnames(x) = c("A", "B","C")
dependence.structure(x) # names of variables are used as labels
dependence.structure(coins(100),vec = c(1,1,2))
# 3-dependent rv of which the first two rv are used together as one rv, thus 2-dependence.
dependence.structure(x,vec = c(1,1,2)) # names of variables are used as labels
dependence.structure(cbind(coins(200), coins(200,k=5)), verbose = TRUE)
#1,2,3 are 3-dependent, 4,..,9 are 6-dependent
# similar to the the previous example, but
# the pair 1,3 is treated as one sample,
# anagously the pair 2,4. In the resulting structure one does not
# see anymore that the dependence of 1,2,3,4 with the rest is due
# to 4.
dependence.structure(cbind(coins(200), coins(200,k=5)),
    vec = c(1, 2, 1, 2, 3, 4, 5, 6, 7),verbose = TRUE)
```

\#\#\# Advanced:

```
# How to check the empirical power of the detection algorithm?
# Use a dataset for which the structure is detected, e.g. dep_struct_several_26_100.
# run:
dependence.structure(dep_struct_several_26_100,
                                    detection.aim = list(c(ncol(dep_struct_several_26_100))))
# The output provides the first detection aim. Now we run the same line with the added
# detection aim
dependence.structure(dep_struct_several_26_100,detection.aim = list(c(3,1, 1, 1, 2, 2, 2, 3, 4,
    5, 6, 7, 8, 8, 8, 9, 9, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 1, 2, 8, 9),
    c(ncol(dep_struct_several_26_100))))
# and get the next detection aim ... thus we finally obtain all detection aims.
# now we can run the code with new sample data ....
N = 100
dependence.structure(cbind(coins(N,2), tetrahedron(N),coins(N,4), tetrahedron(N),
                    tetrahedron(N),coins(N,3),coins(N,3),rnorm(N)),
                    detection.aim = list(c(3,1, 1, 1, 2, 2, 2, 3, 4, 5, 6, 7, 8, 8, 8,
    9, 9, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 1, 2, 8, 9),
    c(4,1, 1, 1, 2, 2, 2, 3, 4, 5, 6, 7, 8, 8, 8, 9, 9, 9, 10, 10, 10, 10, 11, 11, 11,
        11, 12, 1, 2, 8, 9, 10, 11),
    c(5, 1, 1, 1, 2, 2, 2, 3, 3, 3, 3, 3, 4, 4, 4, 5, 5, 5, 6, 6, 6, 6, 7, 7, 7, 7, 8, 1,
        2, 4, 5, 6, 7, 3),
    c(5, 1, 1, 1, 2, 2, 2, 3, 3, 3, 3, 3, 4, 4, 4, 5, 5, 5, 6, 6, 6, 6, 7, 7, 7, 7, 8, 1,
        2, 4, 5, 6, 7, 3)))$detected
# ... and one could start to store the results and compute the rate of successes.
# ... or one could try to check how many samples are necessary for the detection:
re = numeric(100)
for (i in 2:100) {
    re[i] =
        dependence.structure(dep_struct_several_26_100[1:i,],verbose = FALSE,
                            detection.aim = list(c(3,1, 1, 1, 2, 2, 2, 3, 4, 5, 6, 7, 8,
                8, 8, 9, 9, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 1, 2, 8, 9),
            c(4,1, 1, 1, 2, 2, 2, 3, 4, 5, 6, 7, 8, 8, 8, 9, 9, 9, 10, 10, 10, 10, 11, 11,
                    11, 11, 12, 1, 2, 8, 9, 10, 11),
                c(5, 1, 1, 1, 2, 2, 2, 3, 3, 3, 3, 3, 4, 4, 4, 5, 5, 5, 6, 6, 6, 6, 7, 7, 7, 7,
                    8, 1, 2, 4, 5, 6, 7, 3),
                c(5, 1, 1, 1, 2, 2, 2, 3, 3, 3, 3, 3, 4, 4, 4, 5, 5, 5, 6, 6, 6, 6, 7, 7, 7, 7,
                    8, 1, 2, 4, 5, 6, 7, 3)))$detected
    print(paste("First", i,"samples. Detected?", re[i]==1))
}
cat(paste("Given the 1 to k'th row the structure is not detected for k=",which(re == FALSE),"\n"))
```


## Description

It was generated by

```
set.seed(532333356)
N = 100
x = matrix(sample.int(2,10*N,replace = TRUE)-1,ncol = 10)
for (i in c(2,5,9)) x = cbind(x,(rowSums(as.matrix(x[,1:(i-1)]))
dep_struct_iterated_13_100 = x
save(dep_struct_iterated_13_100,file ="dep_struct_iterated_13_100.rda")
```


## Usage

dep_struct_iterated_13_100

## Format

matrix 13 variables (columns), 100 independent samples (rows)

## Details

To avoid irritation, note that the seed is just a simple integer hash value of the variable name.

```
dep_struct_ring_15_100
    example dataset for dependence.structure
```


## Description

It was generated by
set. seed (436646700)
$\mathrm{N}=100$
$\mathrm{n}=15$
$\mathrm{x}=$ matrix(sample.int $(2, \mathrm{~N} * \mathrm{n}$, replace $=$ TRUE) -1 , nrow $=\mathrm{N}$ )
$x[, 4]=\operatorname{rowSums}(x[, 1: 3])$
$x[, 7]=\operatorname{rowSums}(x[, 4: 6])$
$x[, 10]=\operatorname{rowSums}(x[, 7: 9])$
$x[, 13]=\operatorname{rowSums}(x[, 10: 12])$
$x[, 15]=\operatorname{rowSums}(x[, c(13,14,1)])$
dep_struct_ring_15_100 = x
save(dep_struct_ring_15_100,file ="dep_struct_ring_15_100.rda")

## Usage

dep_struct_ring_15_100

## Format

matrix 15 variables (columns), 100 independent samples (rows)

## Details

To avoid irritation, note that the seed is just a simple integer hash value of the variable name.

```
dep_struct_several_26_100
```

    example dataset for dependence.structure
    
## Description

It was generated by
set. seed(1348879148)
$\mathrm{N}=100$
dep_struct_several_26_100 = cbind(coins(N, 2), tetrahedron(N), coins(N,4), tetrahedron ( N ), tetrahedron( N ), coins( $\mathrm{N}, 3$ ), coins(N,3), rnorm(N))
save(dep_struct_several_26_100,file ="dep_struct_several_26_100.rda")

## Usage

dep_struct_several_26_100

## Format

matrix 26 variables (columns), 100 independent samples (rows)

## Details

To avoid irritation, note that the seed is just a simple integer hash value of the variable name.

```
dep_struct_star_9_100 example dataset for dependence.structure
```


## Description

It was generated by
set.seed(222454572)
N = 100
$y=\operatorname{coins}(N, 2)$
dep_struct_star_9_100 = cbind(y,y,y)
save(dep_struct_star_9_100,file ="dep_struct_star_9_100.rda")

## Usage

dep_struct_star_9_100

## Format

matrix 9 variables (columns), 100 independent samples (rows)

## Details

To avoid irritation, note that the seed is just a simple integer hash value of the variable name.

```
emp.transf Monte Carlo empirical transform
```


## Description

Transforms a matrix (rows: samples, columns: variables) into a matrix of uniform samples with the same dependence structure via the Monte Carlo empirical transform.

## Usage

emp.transf(x, continuous = FALSE)

## Arguments

x
data matrix (rows: samples, columns: variables)
continuous boolean, if TRUE it provides the classical (non-Monte-Carlo) transformation by the empirical distribution function, which is a reasonable choice for data of continuous distributions.

## References

For the theoretic background see the reference [5] given on the main help page of this package: multivariance-package.

```
fastdist fast Euclidean distance matrix
```


## Description

fast Euclidean distance matrix

## Usage

fastdist(x)

## Arguments

x
matrix with sample rows for which the distance matrix is computed (to use with vectors, use as.matrix(x))

## Examples

```
#require(microbenchmark)
#x = rnorm(100)
#microbenchmark(fastdist(as.matrix(x)),as.matrix(dist(x)))
```

fastEuclideanCdm fast centered Euclidean distance matrix

## Description

fast centered Euclidean distance matrix

## Usage

fastEuclideanCdm(x, normalize)

## Arguments

$x \quad$ matrix with sample rows for which the distance matrix is computed (to use with vectors, use as.matrix(x))
normalize boolean. If TRUE the matrix will be normalized to mean 1.
find.cluster cluster detection

## Description

Performs the detection of dependence structures algorithm until a cluster is found. This function is the basic building block dependence. structure. Advanced users, might use it directly.

## Usage

find.cluster(
x ,
vec $=1: \operatorname{ncol}(x)$,
list.cdm = cdms(x, vec = vec),
mem = as.numeric(1:max (vec)),
cluster.to.vertex $=1: \max (m e m)$, vertex.to.cdm = 1:max(mem), previous.n.o.cdms = rep(0, max(mem)), all.multivariances $=$ numeric(0),
g = igraph: :add.vertices(igraph::graph.empty(, directed = FALSE), max(mem), label = sapply(1:max(mem), function(r) paste(colnames(x, do.NULL = FALSE, prefix = "")[vec == $r]$, collapse = ",")), shape = "circle"),
fixed.rejection.level = NA,

```
    alpha = 0.05,
    p.adjust.method = "holm",
    verbose = TRUE,
    kvec = 2:max(mem),
    parameter.range = NULL,
    type = "conservative",
    stop.too.many = NULL,
    ..
)
```


## Arguments

| x |
| :--- |
| vec |
| list.cdm |
| mem |$\quad$| vector, it indicates which columns are initially treated together as one sample |
| :--- |
| list of doubly centered distance matrices |
| numeric vector, its length is the number of vertices, its content is the number |
| of the corresponding cluster for the current iteration, i.e., vertex i belongs to |
| cluster mem[i] |

cluster.to.vertex
vector, contains the cluster to vertex relations, i.e., cluster. to.vertex[i] is
the index of the vertex which represents cluster i

```
type the method for the detection, one of 'conservative','resample','pearson_approx'
    or 'consistent'.
stop.too.many numeric, upper limit for the number of tested tuples. A warning is issued if it is
    used. Use stop. too.many = NULL for no limit.
... are passed to resample.multivariance in the case of 'type = resample'
```


## Details

For further details see dependence. structure.

```
independence.test testfor independence
```


## Description

Depreciated. Use multivariance.test instead. It provides all options and returns test result in a standard R format.

## Usage

```
independence.test(
    x,
    vec = 1:ncol(x),
    alpha = 0.05,
    type = "distribution_free",
    verbose = TRUE,
)
```


## Arguments

alpha significance level
x
vec
type
verbose
. . .
either a data matrix or a list of doubly centered distance matrices
if x is a matrix, then this indicates which columns are treated together as one sample; if x is a list, these are the indexes for which the multivariance is calculated. The default is all columns and all indexes, respectively.
one of "pearson_approx", "distribution_free", "resample"
logical, if TRUE meaningful text output is generated.
these are passed to cdms (which is only invoked if $x$ is a matrix)

## Details

This computes a test of independence for the columns of a sample matrix (required for the resampling test) or for given doubly centered distance matrices (only possible for the distribution-free test).

The "pearson_approx" and "resample" are approximately sharp. The latter is based on a resampling approach and thus much slower. The "distribution_free" test might be very conservative. The doubly centered distance matrices can be prepared by cdms. But note that for the test based on Pearson's approximation and for the resampling test, the data matrix has to be given.

## Value

Returns TRUE if the hypothesis of independence is NOT rejected, otherwise FALSE.

## References

For the theoretic background see the references given on the main help page of this package: multivariance-package.

## Examples

```
independence.test(coins(100)) #dependent sample which is 2-independent
independence.test(coins(100),type = "resample") #dependent sample which is 2-independent
independence.test(coins(100)[,2:3]) # independent sample
independence.test(coins(100)[,2:3],type = "resample") # independent sample
independence.test(coins(10), type = "resample") #dependent sample which is 2-independent
independence.test(coins(10)[,2:3],type = "resample") #dependent sample which is 2-independent
```

layout_on_circles special igraph layout for the dependence structure visualization

## Description

It places the variable nodes on an outer circle and the dependency nodes on an inner circle

## Usage

layout_on_circles(g, $\mathrm{n}=$ sum(is.na(igraph:: $\mathrm{V}(\mathrm{g}) \$ 1$ level)))

## Arguments

g
graph
n
number of vertices on outer circle

## Details

This is the standard layout for the full dependence structure, since in this case there often too many nodes which make the other (usual) layout incomprehensible.

## Examples

```
N = 200
y = coins(N,2)
x = cbind(y,y,y)
g = dependence.structure(x,structure.type = "clustered",verbose = FALSE)$graph
plot(g)
plot(g,layout = layout_on_circles(g))
```

m.multivariance $\quad m$ distance multivariance

## Description

Computes m distance multivariance.

## Usage

```
m.multivariance(
    x,
    vec = NA,
    m = 2,
    Nscale = TRUE,
    Escale = TRUE,
    squared = TRUE,
    ...
    )
```


## Arguments

x
vec if $x$ is a matrix, then this indicates which columns are treated together as one sample; if $x$ is a list, these are the indexes for which the multivariance is calculated. The default is all columns and all indexes, respectively.
$\mathrm{m} \quad=2$ or 3 the m-multivariance will be computed.
Nscale if TRUE the multivariance is scaled up by the sample size (and thus it is exactly as required for the test of independence)
Escale if TRUE then it is scaled by the number of multivariances which are theoretically summed up (in the case of independence this yields for normalized distance matrices an estimator with expectation 1)
squared if FALSE it returns the actual multivariance, otherwise the squared multivariance (less computation)
... these are passed to cdms (which is only invoked if x is a matrix)

## Details

$m$-distance multivariance is per definition the scaled sum of certain distance multivariances, and it characterize m-dependence.

As a rough guide to interpret the value of total distance multivariance note:

- Large values indicate dependence.
- If the random variables are (m-1)-independent and Nscale $=$ TRUE, values close to 1 and smaller indicate $m$-independence, larger values indicate dependence. In fact, in the case of independence the test statistic is a Gaussian quadratic form with expectation 1 and samples of it can be generated by resample.multivariance.
- If the random variables are $(\mathrm{m}-1)$-independent and Nscale $=$ FALSE, small values (close to 0 ) indicate $m$-independence, larger values indicate dependence.

Since random variables are always 1 -independent, the case $m=2$ characterizes pairwise independence.

Finally note, that due to numerical (in)precision the value of m-multivariance might become negative. In these cases it is set to 0 . A warning is issued, if the value is negative and further than the usual (used by all. equal) tolerance away from 0 .

## References

For the theoretic background see the reference [3] given on the main help page of this package: multivariance-package.

## Examples

```
x = matrix(rnorm(3*30),ncol = 3)
# the following values are identical
m.multivariance(x,m =2)
1/choose(3,2)*(multivariance(x[,c(1,2)]) +
    multivariance(x[,c(1,3)]) +
    multivariance(x[,c(2,3)]))
# the following values are identical
m.multivariance(x,m=3)
multivariance(x)
# the following values are identical
1/4*(3*(m.multivariance(x,m=2)) + m.multivariance(x,m=3))
total.multivariance(x, Nscale = TRUE)
1/4*(multivariance(x[,c(1,2)], Nscale = TRUE) +
    multivariance(x[,c(1,3)], Nscale = TRUE) +
    multivariance(x[,c(2,3)], Nscale = TRUE) + multivariance(x, Nscale = TRUE))
```

```
multicorrelation distance multicorrelation
```


## Description

Computes various types of sample distance multicorrelation as defined and discussed in $[3,4,6]$.

```
Usage
multicorrelation(
    x,
    vec = 1:ncol(x),
    type = "total.upper.lower",
    multicorrelation.type = "normalized",
    estimator.type = "bias.corrected",
    squared = TRUE,
    ...
)
Mcor(
    x,
    vec = 1:ncol(x),
    type = "total.upper.lower",
    multicorrelation.type = "normalized",
    estimator.type = "bias.corrected",
    squared = TRUE,
)
```


## Arguments

| X | either a data matrix or a list of doubly centered distance matrices |
| :---: | :---: |
| vec | if x is a matrix, then this indicates which columns are treated together as one sample; if $x$ is a list, these are the indexes for which the multivariance is calculated. The default is all columns and all indexes, respectively. |
| type | default: "total.lower.upper", for details and other options see below |
| multicorrelation.type |  |
|  | one of "normalized", "unnormalized" |
| estimator.type | one of "biased", "bias. corrected" |
| squared | if FALSE it returns the actual multivariance, otherwise the squared multivariance (less computation) |
|  | these are passed to cdms (which is only invoked if x is a matrix) |

## Details

There exist many variants of distance multicorrelation as discussed in [6] - and only in specific cases a direct comparison of the values is meaningful.
The implemented options are:

- total.upper.lower normalized bias.corrected: default; bounded by 1 ; fast; population limit characterizes independence by 0
- pairwise normalized bias.corrected: bounded by 1; fast; population limit characterizes pairwise independence by 0
- total. upper normalized biased: biased versions of the above
- total.lower normalized biased
- pairwise normalized biased
- multi normalized biased: population limit characterizes only in case of lower independence the independence of all variables by 0
- m.multi. 3 normalized biased: population limit characterizes only in case of pairwise independence the 3 -independence of all variables by 0
- pairwise unnormalized biased population limit characterizes pairwise independence by 0 and relation by similarity transforms by 1
- multi unnormalized biased: population limit characterizes only in case of lower independence the independence of all variables by 0 and relation by similarity transforms by 1
- m.multi. 3 unnormalized biased: population limit characterizes only in case of pairwise independence the 3 -independence of all variables by 0 and relation by similarity transforms by 1


## Further details:

The "bias. corrected" versions require a data matrix, since they compute bias corrected centered distance matricies.
For "multi" the unnormalized and normalized version coincide if an even number of variables is considered. They usually differ if an odd number of variables is considered. If all variables are related by similarity transforms the unnormalized "unnormalized" multicorrelations are 1.
For "pairwise" an alias is "m.multi.2".
For total multicorrelation there is currently only a feasible empirical estimator for a lower or upper bound. These are upper and lower bounds for in the population setting. When using bias corrected estimators these are in general no proper bounds, but their range can be used as values for comparisons.

## Value

Value of the multicorrelation(s).

## References

For the theoretic background see the references $[2,3,6]$ given on the main help page of this package: multivariance-package.

## Examples

```
y = rnorm(100)
x = cbind(y,y*2,(y-2)/3,y+1,y*5) # all variables are related by similarity transforms
# compute all types of correlations for x:
for (ty in c("total.lower","total.upper","pairwise","m.multi.3","multi"))
    for (mty in c("normalized"))
        print(paste(format(multicorrelation(
        x,type=ty,multicorrelation.type = mty,estimator.type = "biased")
        ,digits=3,nsmall = 3,width = 7),mty,ty,"correlation - biased estimate"))
for (ty in c("total.upper.lower","pairwise"))
    for (mty in c("normalized"))
        print(paste(format(multicorrelation(
        x,type=ty,multicorrelation.type = mty,estimator.type = "bias.corrected")
        ,digits=3,nsmall = 3,width = 7),mty,ty,"correlation - bias corrected estimate"))
for (ty in c("m.multi.2","m.multi.3","multi"))
    for (mty in c("unnormalized"))
        print(paste(format(multicorrelation(
        x,type=ty,multicorrelation.type = mty,estimator.type = "biased")
        ,digits=3,nsmall = 3,width = 7),mty,ty,"correlation - biased estimate"))
```

multivariance distance multivariance

## Description

Computes the distance multivariance, either for given data or a given list of doubly centered distance matrices.

## Usage

multivariance(
x ,
$\mathrm{vec}=\mathrm{NA}$,
Nscale = TRUE,
correlation = FALSE,
squared $=$ TRUE,
...
)

## Arguments

x
either a data matrix or a list of doubly centered distance matrices
vec
if $x$ is a matrix, then this indicates which columns are treated together as one sample; if x is a list, these are the indexes for which the multivariance is calculated. The default is all columns and all indexes, respectively.

| Nscale | if TRUE the multivariance is scaled up by the sample size (and thus it is exactly <br> as required for the test of independence) |
| :--- | :--- |
| correlation | depreciated, please use the function multicorrelation instead. <br> squared |
| if FALSE it returns the actual multivariance, otherwise the squared multivariance <br> (less computation) |  |
| $\ldots$ | these are passed to cdms (which is only invoked if $x$ is a matrix) |

## Details

If $x$ is a matrix and $v e c$ is not given, then each column is treated as a separate sample. Otherwise vec has to have as many elements as $x$ has columns and values starting from 1 up to the number of 'variables', e.g. if $x$ is an $N$ by 5 matrix and vec $=c(1,2,1,3,1)$ then the multivariance of the 1 -dimensional variables represented by column 2 and 4 and the 3 -dimensional variable represented by the columns $1,3,5$ is computed.
As default it computes the normalized Nscaled squared multivariance, for a multivariance without normalization the argument normalize $=$ FALSE has to be passed to cdms.
correlation $=$ TRUE yields values between 0 and 1 . These can be interpreted similarly to classical correlations, see also multicorrelation.
As a rough guide to interpret the value of distance multivariance note:

- If the random variables are not ( $\mathrm{n}-1$ )-independent, large values indicate dependence, but small values are meaningless. Thus in this case use total.multivariance.
- If the random variables are ( $\mathrm{n}-1$ )-independent and Nscale $=$ TRUE, values close to 1 and smaller indicate independence, larger values indicate dependence. In fact, in the case of independence the test statistic is a Gaussian quadratic form with expectation 1 and samples of it can be generated by resample.multivariance.
- If the random variables are $(\mathrm{n}-1)$-independent and Nscale $=$ FALSE, small values (close to 0 ) indicate independence, larger values indicate dependence.

Finally note, that due to numerical (in)precision the value of multivariance might become negative. In these cases it is set to 0 . A warning is issued, if the value is negative and further than the usual (used by all. equal) tolerance away from 0 .

## References

For the theoretic background see the references given on the main help page of this package: multivariance-package.

## Examples

multivariance(matrix(rnorm(100*3), ncol = 3)) \#independent sample multivariance(coins(100)) \#dependent sample which is 2 -independent

```
x = matrix(rnorm(100*2),ncol = 2)
x = cbind(x,x[,2])
multivariance(x) #dependent sample which is not 2-independent (thus small values are meaningless!)
multivariance(x[,1:2]) #these are independent
multivariance(x[,2:3]) #these are dependent
```

```
    multivariance(x[,2:3],correlation = TRUE)
```

multivariance.pvalue transform multivariance to p-value

## Description

Computes a conservative p-value for the hypothesis of independence for a given multivariance / m-multivariance / total multivariance.

## Usage <br> multivariance.pvalue(x)

## Arguments

x
value of a normalized multivariance scaled by the sample size (i.e., computed with Nscale = TRUE)

## Details

This is based on a distribution-free approach. The p-value is conservative, i.e. it might be much smaller. This is the counterpart to rejection. level. For a less conservative approach see resample.pvalue or pearson. pvalue.
p-values larger than 0.215 might be incorrect, since the distribution-free estimate on which the computation is based only holds up to 0.215 .

## References

For the theoretic background see the references given on the main help page of this package: multivariance-package.
multivariance.test independence tests based on (total-/2-/3-) multivariance

## Description

This performs the (specified by type and p.value.type) independence test for the columns of a sample matrix.

```
Usage
    multivariance.test(
        x,
        vec = 1:ncol(x),
        type = "total",
        p.value.type = "pearson_approx",
        verbose = TRUE,
    )
```


## Arguments

x
vec
type
p.value.type

## verbose

... these are passed to cdm

## Details

For the use of vec see the examples below and the more detailed explanation of this argument for multivariance.
The types "independence" and "total" are identical: an independence test is performed.
Also the types "pairwise independence" and "m.multi. 2 " are identical: a test of pairwise independence is performed.
The type "m.multi. 3 ", performs a test for 3-independence, assuming pairwise independence. The type "multi" performs a test for $n$-independence, assuming ( $\mathrm{n}-1$ )-independence.
There are several ways (determined by p.value. type) to estimate the p-value: The "pearson_approx" and "resample" are approximately sharp. The latter is based on a resampling approach and thus much slower. The "distribution_free" test might be very conservative, its p-value estimates are only valid for p -values lower than 0.215 - values above should be interpreted as "values larger than 0.215 ". Finally, "pearson_unif" uses fixed parameters in Pearson's estimate, it is only applicable for univariate uniformly distributed marginals
All tests are performed using the standard euclidean distance. Other distances can be supplied via the . . ., see cdm for the accepted arguments.

## Value

A list with class "htest" containing the following components:
statistic the value of the test statistic,
$p$.value the $p$-value of the test statistic,
method a character string indicating the type of test performed,
data. name a character string giving the name(s) of the data.

## References

For the theoretic background see the references given on the main help page of this package: multivariance-package.

## Examples

```
# an independence test
multivariance.test(dep_struct_several_26_100,p.value.type = "distribution_free") # conservative
multivariance.test(dep_struct_several_26_100,p.value.type = "resample") #sharp but slow
multivariance.test(dep_struct_several_26_100,p.value.type = "pearson_approx") #
# as an example, all tests for one data set:
coins100 = coins(100)
for (ty in c("total","m.multi.2","m.multi.3","multi"))
    for (pvt in c("distribution_free","resample","pearson_approx"))
        print(multivariance.test(coins100,type=ty,p.value.type = pvt))
# using the vec argument:
x = matrix(rnorm(50*6),ncol = 10) # a 50x6 data matrix
vec = c(1, 2,3,4,5,6) # each column is treated as one variable
multivariance.test(x,vec,p.value.type = "distribution_free") # is the same as the default
vec =c(1, 2, 2, 1, 3, 1)
# column 1,4,6 are treated as one variable
# column 2,3 are treated as one variable
# column 5 is treated as one variable
multivariance.test(x,vec,p.value.type = "distribution_free")
```

```
multivariance.timing estimate of the computation time
```


## Description

Estimates the computation time. This is relative rough. First run with determine.parameters $=$ TRUE (which takes a while). Then use the computed parameters to determine the computation time/or sample size.

```
Usage
    multivariance.timing(
    N = NULL,
    n,
    sectime = NULL,
    coef.cdm = 15.2,
    coef.prod = 2.1,
    coef.sum = 1.05,
    determine.parameters = FALSE
)
```


## Arguments

N
number of samples. If NULL and sectime is given, then $N$ is computed.
n
number of variables
sectime desired computation time in seconds. If NULL then the required computation time is computed.
coef.cdm computation time parameter for the doubly centered distance matrices
coef.prod computation time parameter for matrix products
coef.sum computation time parameter for matrix sums
determine.parameters
if TRUE then the parameters for the current computer are determined. This might take a while (3 loops to $\mathrm{N}=1000$ ).

## Details

When detecting the parameters, the median of the computation times is used.

## Examples

```
Ns = (1:100)*10
ns = 1:100
fulltime = outer(Ns,ns,FUN = function(N,n) multivariance.timing(N,n))
contour(Ns,ns,fulltime,xlab = "N",ylab = "n",
    main = "computation time of multivariance in secs",
    sub = "using default parameters -
    use 'determine.parameters = TRUE' to compute machine specific values")
# Run to determine the parameters of your system:
# multivariance.timing(determine.parameters = TRUE)
```

multivariances.all simultaneous computation of multivariance and total/ 2-/ 3-
multivariance

## Description

Computes simultaneously multivariance, total multivariance, 2-multivariance and 3-multivariance.

## Usage

multivariances.all(x, vec $=$ NA, Nscale $=$ TRUE, squared $=$ TRUE, ...)

## Arguments

x
vec

Nscale if TRUE the multivariance is scaled up by the sample size (and thus it is exactly as required for the test of independence)
squared if FALSE it returns the actual multivariance, otherwise the squared multivariance (less computation)
... these are passed to cdms (which is only invoked if $x$ is a matrix)

## Details

The computation is faster than the separate computations.

## Value

Returns a vector with multivariance, total.multivariance, 2-multivariance and 3-multivariance

```
See Also
multivariance, total.multivariance, m.multivariance
```


## Examples

```
x = coins(100,k = 3)
multivariances.all(x)
# yields the same as:
multivariance(x)
total.multivariance(x)
m.multivariance(x,m=2)
m.multivariance(x,m=3)
```

pearson.pvalue fast p-value approximation

## Description

Computes the p-value of a sample using Pearson's approximation of Gaussian quadratic forms with the estimators developed by Berschneider and Böttcher in [4].

## Usage

pearson.pvalue (x, vec = NA, type = "multi", ...)

## Arguments

| $x$ | matrix, the rows should be iid samples |
| :--- | :--- |
| vec | vector, which indicates which columns of $x$ are treated together as one sample. |
|  | The default case treats each column as a separate sample. |
| type | one of "multi", "total", "m.multi. $2 ", " m . m u l t i .3 ", " a l l " ~$ |
| $\ldots$. | these are passed to cdms |

## Details

This is the method recommended in [4], i.e., using Pearson's quadratic form estimate with the unbiased finite sample estimators for the mean and variance of normalized multivariance together with the unbiased estimator for the limit skewness.

## References

For the theoretic background see the reference [4] given on the main help page of this package: multivariance-package.

```
pearson.qf approximate distribution function of a Gaussian quadratic form
```


## Description

Approximation of the of the value of the distribution function of a Gaussian quadratic form based on its first three moments.

## Usage

pearson. $\mathrm{qf}(\mathrm{x}$, moment, lower.tail $=$ TRUE, verbose $=$ FALSE)

## Arguments

X
moment
lower.tail
verbose
value at which the distribution function is to be evaluated vector with the mean, variance and skewness of the quadratic form logical, indicating of the lower or upper tail of the distribution function should be calculated
logical, if TRUE a warning is issued if negative moments are sanitized to 0 .

## Details

This is Pearson's approximation for Gaussian quadratic forms as stated in [4] (equation (4.65) in arXiv:1808.07280v2)

## References

For the theoretic background see the reference [4] given on the main help page of this package: multivariance-package.

## Description

Under independence the probability for the normalized and Nscaled (squared) multivariance to be above this level is less than alpha. The same holds for the normalized, Nscaled and Escaled (squared) total multivariance and m-multivariance.

## Usage

rejection.level(alpha)

## Arguments

> alpha level of significance

## Details

This is based on a distribution-free approach. The value might be very conservative. This is the counterpart to multivariance.pvalue. For a less conservative approach see resample. rejection.level.

The estimate is only valid for alpha smaller than 0.215 .

## Examples

```
rejection.level(0.05) #the rejection level, for comparison with the following values
total.multivariance(matrix(rnorm(100*3),ncol = 3)) #independent sample
total.multivariance(coins(100)) #dependent sample which is 2-independent
# and the p values are (to compare with alpha)
multivariance.pvalue(total.multivariance(matrix(rnorm(100*3),ncol = 3))) #independent sample
multivariance.pvalue(total.multivariance(coins(100))) #dependent sample which is 2-independent
## Not run:
# visualization of the rejection level
curve(rejection.level(x),xlim = c(0.001,0.215),xlab = "alpha")
## End(Not run)
```

```
resample.multivariance
```

    resampling (total/m-) multivariance
    
## Description

The distribution of the test statistic under the hypothesis of independence is required for the independence tests. This function generates approximate samples of this distribution either by sampling without replacement (permutations) or by sampling with replacement (bootstrap).

## Usage

resample.multivariance(
x ,
vec = 1:ncol(x),
times $=300$,
type = "multi",
resample.type = "permutation",
)

## Arguments

X
vec vector, which indicates which columns of $x$ are treated together as one sample
times integer, number of samples to generate
type one of "multi","total","m.multi.2","m.multi.3","all"
resample.type one of "permutation", "bootstrap". The samples are generated without replacement (permutations) or with replacement (bootstrap).
... is passed to cdms, multivariance, total.multivariance, m.multivariance, respectively.

## Details

The resampling is done by sampling from the original data either without replacement ("permutation") or with replacement ("bootstrap"). Using resampling without replacement is (much) faster (due to special identities which only hold in this case).
For convenience also the actual (total /m-) multivariance is computed and its p-value.

## Value

A list with elements
resampled the (total/m-)multivariances of the resampled data,
original the (total/m-)multivariance of the original data,
p . value the p -value of the original data, computed using the resampled data

## References

For the theoretic background see the reference [3] given on the main help page of this package: multivariance-package.

## Examples

```
re.m = resample.multivariance(matrix(rnorm(30*2),nrow = 30),
    type= "multi",times = 300)$resampled
curve(ecdf(re.m)(x), xlim = c(0,4),main = "empirical distribution of the test statistic under H_0")
```

    resample.pvalue \(\quad p\)-value via resampling
    
## Description

Use a resampling method to generate samples of the test statistic under the hypothesis of independence. Based on these the p.value of a given value of a test statistic is computed.

## Usage

resample.pvalue(value, ...)

## Arguments

value numeric, the value of (total-/m-)multivariance for which the p -value shall be computed
... passed to resample.multivariance. Required is the data matrix x.

## Details

This function is useful if a $p$-value of a test statistic shall be computed based on the resampling values of the test statistic of a different sample. For the p-value based on the same sample resample.multivariance(...)\$p.v. is sufficient.

## Value

It returns 1 minus the value of the empirical distribution function of the resampling samples evaluated at the given value.

## References

For the theoretic background see the reference [3] given on the main help page of this package: multivariance-package.

## Examples

```
\(x=\) coins(100)
resample.pvalue(multivariance (x), \(x=x\),times \(=300\) )
resample.pvalue(multivariances.all(x), \(x=x\), times \(=300\),type \(=\) "all")
```

resample.rejection.level
rejection level via resampling

## Description

Uses the resample method to sample from the test statistic under the hypothesis of independence. The alpha quantile of these samples is returned.

## Usage

resample.rejection.level(alpha = 0.05, ...)

## Arguments

| alpha | numeric, the significance value |
| :--- | :--- |
| $\ldots$ | passed to resample.multivariance. Required is the data matrix $x$. |

## References

For the theoretic background see the reference [3] given on the main help page of this package: multivariance-package.

## Examples

```
resample.rejection.level(0.05,matrix(rnorm(30*2),nrow = 30))
resample.rejection.level(0.05,matrix(rnorm(30*3),nrow = 30),vec = c(1,1,2))
```


## Description

resamples doubly centered distance matrices

## Usage

sample.cdms(list.cdm, replace $=$ FALSE, incl.first $=$ FALSE)

## Arguments

| list.cdm | a list of doubly centered distance matrices |
| :--- | :--- |
| replace | boolean, sampling with or without replacement |
| incl.first | boolean, if TRUE also the first component is resampled |

## Value

Returns a list of doubly centered distance matrices, each matrix corresponds to the resampled columns of the corresponding sample, using resampling with replacement (bootstrap) or without replacement (permutations).

## Description

resample the columns of a matrix

## Usage

sample.cols(x, vec $=1: n c o l(x)$, replace $=$ TRUE, incl.first $=$ TRUE)

## Arguments

x
matrix
vec vector, indicates which columns belong together
replace boolean, sampling with or without replacement
incl.first boolean, if TRUE also the first component is resampled

## Value

Returns a matrix with the same dimensions as x . The columns are resampled from the original columns. The resampling is done with replacement (replace $=$ TRUE) or without (replace $=$ FALSE). Columns which belong together (indicated by vec) are resampled identically, i.e., all values in rows of these are kept together.

## Examples

```
sample.cols(matrix(1:15, nrow = 5),vec = c(1,1,2))
```


## Description

This function creates samples of a tetrahedron-dice colored $\mathrm{r}, \mathrm{g}, \mathrm{b}$ and rgb . Each sample indicates if for the thrown dice the colors $\mathrm{r}, \mathrm{g}$ and b are contained on the bottom side of the dice.

## Usage

tetrahedron( $\mathrm{N}=1000$ )

## Arguments

$N \quad$ number of samples

## Value

It returns the samples of the events $\mathrm{r}, \mathrm{g}$ and b as rows of $\mathrm{a} N$ by 3 matrix (the first column corresponds to r , the second to $\mathrm{g}, \ldots$ ). TRUE indicates that this color is on the bottom side of the dice. The columns are dependent but 2-independent.

## References

For the theoretic background see the reference [3] given on the main help page of this package: multivariance-package.

## Examples

tetrahedron(10)

```
    total.multivariance total distance multivariance
```


## Description

computes the total distance multivariance

## Usage

```
total.multivariance(
    x ,
    vec \(=N A\),
    lambda = 1,
    Nscale = TRUE,
    Escale = TRUE,
    squared \(=\) TRUE,
    . . .
)
```


## Arguments

| X | either a data matrix or a list of doubly centered distance matrices |
| :---: | :---: |
| vec | if $x$ is a matrix, then this indicates which columns are treated together as one sample; if $x$ is a list, these are the indexes for which the multivariance is calculated. The default is all columns and all indexes, respectively. |
| lambda | a scaling parameter $>0$. Each k-tuple multivariance gets weight lambda^ $(\mathrm{n}-\mathrm{k})$. |
| Nscale | if TRUE the multivariance is scaled up by the sample size (and thus it is exactly as required for the test of independence) |
| Escale | if TRUE then it is scaled by the number of multivariances which are theoretically summed up (in the case of independence this yields for normalized distance matrices an estimator with expectation 1) |
| squared | if FALSE it returns the actual multivariance, otherwise the squared multivariance (less computation) |
|  | these are passed to cdms (which is only invoked if x is a matrix) |

## Details

Total distance multivariance is per definition the scaled sum of certain distance multivariances, and it characterize dependence.
As a rough guide to interpret the value of total distance multivariance note:

- Large values indicate dependence.
- For Nscale = TRUE values close to 1 and smaller indicate independence, larger values indicate dependence. In fact, in the case of independence the test statistic is a Gaussian quadratic form with expectation 1 and samples of it can be generated by resample.multivariance.
- For Nscale $=$ FALSE small values (close to 0 ) indicate independence, larger values indicate dependence.

Finally note, that due to numerical (in)precision the value of total multivariance might become negative. In these cases it is set to 0 . A warning is issued, if the value is negative and further than the usual (used by all. equal) tolerance away from 0 .

## References

For the theoretic background see the references given on the main help page of this package: multivariance-package.

## Examples

$x=$ matrix (rnorm $(100 * 3)$, ncol $=3$ )
total.multivariance(x) \#for an independent sample
\# the value coincides with
(multivariance(x[,c(1,2)],Nscale = TRUE) + multivariance(x[,c(1,3)],Nscale = TRUE)+ multivariance $(x[, c(2,3)], N s c a l e=T R U E)+$ multivariance $(x, N s c a l e=T R U E)) / 4$
total.multivariance(coins(100)) \#value for a dependent sample which is 2-independent

## Index

```
* datasets
    anscombe.extended, 5
    dep_struct_iterated_13_100,15
    dep_struct_ring_15_100,16
    dep_struct_several_26_100,17
    dep_struct_star_9_100, 17
all.equal, 24, 28, 41
anscombe.extended, 4, 5
cdm, 4, 6, 8, 11, 30
cdms, 4, 7, 11, 21-23, 25, 28, 33, 34, 36, 41
clean.graph, 8,13
CMcor (copula.multicorrelation), 10
coins, 4, }
copula.multicorrelation, 10
copula.multicorrelation.test, 11
copula.multivariance, 3, 11
dep_struct_iterated_13_100, 4, 15
dep_struct_ring_15_100, 4, 16
dep_struct_several_26_100,4,17
dep_struct_star_9_100, 4, 17
dependence.structure, 3, 4, 8, 12,15-17,
    19,21
emp.transf, 4, 18
fastdist,18
fastEuclideanCdm,19
find.cluster, 4, 13,19
independence.test, 21
layout_on_circles,22
m.multivariance, 3, 23, 33, 36
Mcor (multicorrelation), 25
multicorrelation, 3, 10, 25, 28
multivariance, 3, 27, 29, 30, 33, 36
```


## * datasets

```
anscombe.extended, 5
dep_struct_iterated_13_100, 15
dep_struct_ring_15_100, 16
dep_struct_several_26_100, 17
dep_struct_star_9_100, 17
all.equal, 24, 28, 41
anscombe.extended, 4, 5
cdm, 4, 6, \(8,11,30\)
cdms, \(4,7,11,21-23,25,28,33,34,36,41\)
clean.graph, 8,13
CMcor (copula.multicorrelation), 10
coins, 4, 9
copula.multicorrelation, 10
copula.multicorrelation.test, 11
copula.multivariance, 3, 11
dep_struct_iterated_13_100, 4, 15
dep_struct_ring_15_100, 4, 16
dep_struct_several_26_100, 4, 17
dep_struct_star_9_100, 4, 17
dependence.structure, \(3,4,8,12,15-17\), 19, 21
emp.transf, 4, 18
fastdist, 18
find.cluster, 4, 13, 19
independence.test, 21
layout_on_circles, 22
m.multivariance, 3, 23, 33, 36
Mcor (multicorrelation), 25
multivariance, 3, 27, 29, 30, 33, 36
```

multivariance-package, $3,5,7,10-12,14$, $18,22,24,26,28,29,31,34,37,38$, 40, 41
multivariance.pvalue, 3, 7, 29, 35
multivariance.test, 3, 4, 21, 29
multivariance.timing, 4, 31
multivariances.all, 3, 32
p.adjust, 20
p.adjust.methods, 13
pearson.pvalue, 4, 29, 33
pearson.qf, 34
rejection.level, 3, 7, 29, 35
resample.multivariance, 4, 21, 24, 28, 36, 37, 38, 41
resample.pvalue, 4, 29, 37
resample.rejection.level, 4, 35, 38
sample.cdms, 38
sample.cols, 39
tetrahedron, 4, 40
total.multivariance, 3, 28, 33, 36, 40

