

Package: SuperRanker (via r-universe)

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Type Package

Title Sequential Rank Agreement

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Description Tools for analysing the agreement of two or more rankings of the same items. Examples are importance rankings of predictor variables and risk predictions of subjects. Benchmarks for agreement are computed based on random permutation and bootstrap. See Ekstrøm CT, Gerds TA, Jensen, AK (2018). "Sequential rank agreement methods for comparison of ranked lists." *Biostatistics*, 20(4), 582-598 <[doi:10.1093/biostatistics/kxy017](https://doi.org/10.1093/biostatistics/kxy017)> for more information.

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average_overlap	<i>Compute the average overlap</i>
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Description

Compute the average overlap

Usage

```
average_overlap(obj)
```

Arguments

obj Either a vector or matrix

Value

A vector of the average overlap

Examples

```
# setting with 3 lists
mlist <- matrix(cbind(1:8,c(1,2,3,5,6,7,4,8),c(1,5,3,4,2,8,7,6)),ncol=3)
average_overlap(mlist)
```

overlap	<i>Compute the overlap between k ranked lists</i>
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Description

Computes the overlap (number of items present in all k lists divided by the current rank) for each rank in the k lists

Usage

```
overlap(rankMat)
```

Arguments

rankMat	A matrix with k columns corresponding to the k ranked lists. Elements of each column are integers between 1 and the length of the lists
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Value

A vector of the same length as the rows in rankMat containing the overlap between the lists for each rank

Author(s)

Claus Ekstrøm <ekstrom@sund.ku.dk>

plot.sra	<i>Plot sequential rank agreement</i>
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Description

Plot the agreement between lists as a function of the list depth

Usage

```
## S3 method for class 'sra'  
plot(  
  x,  
  xlim,  
  ylim,  
  xlab = "List depth",  
  ylab = "Sequential rank agreement",  
  add = FALSE,  
  ...  
)
```

Arguments

x	Agreement object
xlim	x-axis limits
ylim	y-axis limits
xlab	x-axis lab
ylab	y-axis lab
add	Logical. If TRUE add graph to existing plot.
...	Processed by function <code>prodlim::SmartControl</code> .

Value

Graph

Author(s)

Thomas A. Gerds <tag@biostat.ku.dk>

Examples

```
R1=c(1,2,3,4,5,7,6,8,9,10,11,12,13)
R2=c(5,11,4,7,8,3,12,13,6,10,9,2,1)
a <- sra(list(R1,R2))
plot(a)
arand = colMeans(do.call("rbind",lapply(1:20,function(b){
  sra(list(sample(R1),sample(R1)))
})))
lines(1:length(R1),arand,col=2,lwd=3)

l <- c(1,2,3,4,5,7,6,8,9,10,11,12,13)
l <- 1:100
aa <- sapply(1:20,function(i){
  sra(list(sample(l),sample(l),sample(l)))[i]
})
c(mean(aa),sd(aa))
```

random_list_sra

Simulate sequential rank agreement for randomized unrelated lists

Description

Simulate sequential rank agreement from completely uninformative lists (ie., raw permutations of items) and compute the corresponding sequential rank agreement curves. The following attributes are copied from the input object: number of lists, number of items and amount of censoring.

Usage

```
random_list_sra(
  object,
  B = 1,
  n = 1,
  na.strings = NULL,
  nitems = NULL,
  type = c("sd", "mad"),
  epsilon = 0
)
```

Arguments

object	A matrix of numbers or list of vectors representing ranked lists.
B	An integer giving the number of randomizations to sample over in the case of censored observations
n	Integer: the number of permutation runs. For each permutation run we permute each of the lists in object and compute corresponding the sequential rank agreement curves
na.strings	A vector of character values that represent censored observations
nitems	The total number of items in the original lists if we only have partial lists available. Will be derived from the unique elements of the object if set to NULL (the default)
type	The type of measure to use. Either sd (standard deviation - the default) or mad (median absolute deviance)
epsilon	A non-negative numeric vector that contains the minimum limit in proportion of lists that must show the item. Defaults to 0. If a single number is provided then the value will be recycled to the number of items. Should usually be low.

Value

A matrix with n columns and the same number of rows as for the input object. Each column contains one simulated sequential rank agreement curve from one permutation run.

Author(s)

Claus Ekstrøm <ekstrom@sund.ku.dk>

Examples

```
# setting with 3 lists
m1ist <- matrix(cbind(1:8,c(1,2,3,5,6,7,4,8),c(1,5,3,4,2,8,7,6)),ncol=3)
# compute sequential rank agreement of lists
sra(m1ist)
# compute sequential rank agreement of 5 random permutations
random_list_sra(m1ist, n=5)
```

`smooth_sra`*Smooth quantiles of a matrix of sequential ranked agreements.*

Description

Smooth quantiles of a matrix of sequential ranked agreements.

Usage

```
smooth_sra(object, confidence = 0.95)
```

Arguments

<code>object</code>	A matrix
<code>confidence</code>	the limits to compute

Value

A list containing two vectors for the smoothed lower and upper limits

Author(s)

Claus Ekstrøm <ekstrom@sund.ku.dk>

Examples

```
# setting with 3 lists
mlist <- matrix(cbind(1:8,c(1,2,3,5,6,7,4,8),c(1,5,3,4,2,8,7,6)),ncol=3)
# compute rank agreement of 5 random permutations
null=random_list_sra(mlist,n=15)
# now extract point-wise quantiles according to confidence level
smooth_sra(null)
```

`sra`*Compute the sequential rank agreement*

Description

Compute the sequential rank agreement

Usage

```

sra(object, B, na.strings, nitems, type, epsilon = 0, ...)

## Default S3 method:
sra(object, B, na.strings, nitems, type, epsilon = 0, ...)

## S3 method for class 'matrix'
sra(
  object,
  B = 1,
  na.strings = NULL,
  nitems = nrow(object),
  type = c("sd", "mad"),
  epsilon = 0,
  ...
)

## S3 method for class 'list'
sra(
  object,
  B = 1,
  na.strings = NULL,
  nitems = max(sapply(object, length)),
  type = c("sd", "mad"),
  epsilon = 0,
  ...
)

```

Arguments

<code>object</code>	Either matrix where each column is a ranked list of items or a list of ranked lists of items. Elements are integers between 1 and the length of the lists. The lists should have the same length but censoring can be used by setting the list to zero from a point onwards. See details for more information.
<code>B</code>	An integer giving the number of randomization to sample over in the case of censored observations
<code>na.strings</code>	A vector of strings/values that represent missing values in addition to NA. Defaults to NULL which means only NA are censored values.
<code>nitems</code>	The total number of items in the original lists if we only have partial lists available.
<code>type</code>	The type of measure to use. Either sd (standard deviation - the default) or mad (median absolute deviance around the median)
<code>epsilon</code>	A non-negative numeric vector that contains the minimum limit in proportion of lists that must show the item. Defaults to 0. If a single number is provided then the value will be recycled to the number of items.
<code>...</code>	Arguments passed to methods.

Value

A vector of the sequential rank agreement

Author(s)

Claus Ekstrøm <ekstrom@sund.ku.dk> and Thomas A Gerds <tag@biostat.ku.dk>

Examples

```
m1list <- matrix(cbind(1:8,c(1,2,3,5,6,7,4,8),c(1,5,3,4,2,8,7,6)),ncol=3)
sra(m1list)

m2list <- matrix(cbind(1:8,c(1,2,3,5,6,7,4,8),c(1,5,3,4,2,8,7,6)),ncol=3)
sra(m2list, nitems=20, B=10)

alist <- list(a=1:8,b=sample(1:8),c=sample(1:8))
sra(alist)

blist <- list(x1=letters,x2=sample(letters),x3=sample(letters))
sra(blist)

## censored lists are either too short
clist <- list(x1=c("a","b","c","d","e","f","g","h"),
             x2=c("h","c","f","g","b"),
             x3=c("d","e","a"))
set.seed(17)
sra(clist,na.strings="z",B=10)

## or use a special code for missing elements
Clist <- list(x1=c("a","b","c","d","e","f","g","h"),
             x2=c("h","c","f","g","b","z","z","z"),
             x3=c("d","e","a","z","z","z","z","z"))
set.seed(17)
sra(Clist,na.strings="z",B=10)
```

 sracpp

Compute the sequential rank agreement between k ranked lists

Description

Computes the sequential rank agreement (number of items present in all k lists divided by the current rank) for each rank in the k lists

Usage

```
sracpp(rankMat, maxlength, B, cens, type = 0L, epsilon = as.numeric(c(0)))
```


Arguments

rankMat	A matrix with k columns corresponding to the k ranked lists. Elements of each column are integers between 1 and the length of the lists
maxlength	The maximum depth that are needed XXX
B	The number of resamples to use in the presence of censored lists
cens	A vector of integer values that
type	The type of distance measure to use: 0 (the default) is the variance while 1 is MAD (median absolute deviation)
epsilon	A non-negative numeric vector that contains the minimum limit in proportion of lists that must show the item. Defaults to 0. If a single number is provided then the value will be recycled to the number of items.

Value

A vector of the same length as the rows in rankMat containing the squared (!) sequential rank agreement between the lists for each depth. If the MAD type was chosen then the sequential MAD values are returned

Author(s)

Claus Ekstrøm <ekstrom@sund.ku.dk>

 sracppfull

Compute the sequential rank agreement between k ranked lists

Description

Computes the sequential rank agreement (number of items present in all k lists divided by the current rank) for each rank in the k lists

Usage

```
sracppfull(rankMat, type = 0L, epsilon = as.numeric(c(0)))
```

Arguments

rankMat	A matrix with k columns corresponding to the k ranked lists. Elements of each column are integers between 1 and the length of the lists
type	The type of distance measure to use: 0 (the default) is the variance while 1 is MAD (mean absolute deviation)
epsilon	A non-negative numeric vector that contains the minimum limit in proportion of lists that must show the item. Defaults to 0. If a single number is provided then the value will be recycled to the number of items.

Value

A vector of the same length as the rows in rankMat containing the sequential rank agreement between the lists for each depth (squared for type=0)

Author(s)

Claus Ekstrøm <ekstrom@sund.ku.dk>

SuperRanker

Functions related to comparison of ranked lists

Description

SuperRanker allows you to estimate the agreement between two or more rankings of the same items.

test_sra

Compute a Kolmogorov-Smirnoff-like test for Smooth quantiles of a matrix of sequential rank agreements

Description

Compute a Kolmogorov-Smirnoff-like test for Smooth quantiles of a matrix of sequential rank agreements

Usage

```
test_sra(object, nullobject, weights = 1)
```

Arguments

object	An object created with sra.
nullobject	An object created with random_list_sra.
weights	Either a single value or a vector of the same length as the number of item with the weight that should be given to specific depths.

Value

A single value corresponding to the p-value

Author(s)

Claus Ekstrøm <ekstrom@sund.ku.dk>

Examples

```
# setting with 3 lists
mlist <- matrix(cbind(1:8,c(1,2,3,5,6,7,4,8),c(1,5,3,4,2,8,7,6)),ncol=3)
# compute sequential rank agreements
x=sra(mlist)
# compute rank agreement of 5 random permutations
null=random_list_sra(mlist,n=15)
# now extract point-wise quantiles according to confidence level
test_sra(x,null)
# compare to when we use the result of the first permutation run
test_sra(null[,1],null[,-1])
```

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