

Package ‘idr’

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Type Package

Title Irreproducible Discovery Rate

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Description This is a package for estimating the copula mixture model and plotting correspondence curves. Details are in ``Measuring reproducibility of high-throughput experiments" (2011), Annals of Applied Statistics, Vol. 5, No. 3, 1752-1779, by Li, Brown, Huang, and Bickel.

License GPL (>= 2.0)

Imports stats

LazyLoad yes

NeedsCompilation no

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idr-package

*Irreproducible discovery rate***Description**

This package estimates the reproducibility of observations on a pair of replicate rank lists. It consists of three components: (1) plotting the correspondence curve to visualize reproducibility, (2) quantifying reproducibility using a copula mixture model and estimating the posterior probability for each observation to be irreproducible (local irreproducible discovery rate), and (3) ranking and selecting observations by their irreproducibility.

Details

Package:	idr
Type:	Package
Version:	1.2
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License:	GPL-2
LazyLoad:	yes

The main functions are `est.IDR()`, `get.correspondence()` and `select.IDR()`. `est.IDR` estimates the copula mixture model and the posterior probability for each observation to be irreproducible. `get.correspondence` generates the values for plotting the correspondence curve. `select.IDR` ranks observations by their reproducibility and reports the number of observations passing the specified IDR thresholds.

Author(s)

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References

Q. Li, J. B. Brown, H. Huang and P. J. Bickel. (2011) Measuring reproducibility of high-throughput experiments. *Annals of Applied Statistics*, Vol. 5, No. 3, 1752-1779.

Examples

```
data("simu.idr")
x <- cbind(-simu.idr$x, -simu.idr$y)

mu <- 2.6
sigma <- 1.3
rho <- 0.8
p <- 0.7
```

```
idr.out <- est.IDR(x, mu, sigma, rho, p, eps=0.001, max.ite=20)

names(idr.out)
```

est.IDR	<i>Estimate the irreproducible discovery rate using the copula mixture model</i>
---------	--

Description

Fit a Gaussian copula mixture model.

Usage

```
est.IDR(x, mu, sigma, rho, p, eps=0.001, max.ite=30)
```

Arguments

x	an n by m numeric matrix, where m= num of replicates, n=num of observations. Numerical values representing the significance of the observations. Note that significant signals are expected to have large values of x. In case that smaller values represent higher significance (e.g. p-value), a monotonic transformation needs to be applied to reverse the order before using this function, for example, -log(p-value). Currently, m=2.
mu	a starting value for the mean of the reproducible component.
sigma	a starting value for the standard deviation of the reproducible component.
rho	a starting value for the correlation coefficient of the reproducible component.
p	a starting value for the proportion of reproducible component.
eps	Stopping criterion. Iterations stop when the increment of log-likelihood is < eps*log-likelihood, Default=0.001.
max.ite	Maximum number of iterations. Default=30.

Value

para	estimated parameters: p, rho, mu, sigma.
idr	a numeric vector of the local idr for each observation (i.e. estimated conditional probability for each observation to belong to the irreproducible component).
IDR	a numerical vector of the expected irreproducible discovery rate for observations that are as irreproducible or more irreproducible than the given observations.
loglik	log-likelihood at the end of iterations.
loglik.trace	trajectory of log-likelihood.

Author(s)

Qunhua Li

References

Q. Li, J. B. Brown, H. Huang and P. J. Bickel. (2011) Measuring reproducibility of high-throughput experiments. *Annals of Applied Statistics*, Vol. 5, No. 3, 1752-1779.

Examples

```
data("simu.idr")

# simu.idr$x and simu.idr$y are p-values
# Transfer them such that large values represent significant ones
x <- cbind(-simu.idr$x, -simu.idr$y)

mu <- 2.6
sigma <- 1.3
rho <- 0.8
p <- 0.7

idr.out <- est.IDR(x, mu, sigma, rho, p, eps=0.001, max.ite=20)

names(idr.out)
```

get.correspondence	<i>Compute correspondence profiles</i>
--------------------	--

Description

Compute the correspondence profiles (Psi and Psi') and the corresponding smoothed curve using spline

Usage

```
get.correspondence(x1, x2, t, spline.df = NULL)
```

Arguments

x1	Data values or ranks of the data values on list 1, a vector of numeric values. Large values need to be significant signals. If small values represent significant signals, rank the signals reversely (e.g. by ranking negative values) and use the rank as x1.
x2	Data values or ranks of the data values on list 2, a vector of numeric values. Large values need to be significant signals. If small values represent significant signals, rank the signals reversely (e.g. by ranking negative values) and use the rank as x1.
t	A numeric vector between 0 and 1 in ascending order. t is the right-tail percentage.
spline.df	Degree of freedom for spline, to control the smoothness of the smoothed curve.

Value

psi	the correspondence profile in terms of the scale of percentage, i.e. between (0, 1)
dpsi	the derivative of the correspondence profile in terms of the scale of percentage, i.e. between (0, 1)
psi.n	the correspondence profile in terms of the scale of the number of observations
dpsi.n	the derivative of the correspondence profile in terms of the scale of the number of observations

Each object above is a list consisting of the following items: t: upper percentage (for psi and dpsi) or number of top ranked observations (for psi.n and dpsi.n) value: psi or dpsi smoothed.line: smoothing spline ntotal: the number of observations jump.point: the index of the vector of t such that psi(t[jump.point]) jumps up due to ties at the low values. This only happens when data consists of a large number of discrete values, e.g. values imputed for observations appearing on only one replicate.

Author(s)

Qunhua Li

References

Q. Li, J. B. Brown, H. Huang and P. J. Bickel. (2011) Measuring reproducibility of high-throughput experiments. *Annals of Applied Statistics*, Vol. 5, No. 3, 1752-1779.

Examples

```
# salmon data
data(salmon)

# get.correspondence() needs the observations with high ranks have
# high correlation and the observations with low ranks have low correlation.
# In this dataset, small values have high correlation and large values
# have low correlation.
# Ranking negative values makes the data follow the structure required
# by get.correspondence().
# There are 28 observations in this data set.

rank.x <- rank(-salmon$spawners)
rank.y <- rank(-salmon$recruits)
uv <- get.correspondence(rank.x, rank.y, seq(0.01, 0.99, by=1/28))

# plot correspondence curve on the scale of percentage
plot(uv$psi$t, uv$psi$value, xlab="t", ylab="psi", xlim=c(0, max(uv$psi$t)),
      ylim=c(0, max(uv$psi$value)), cex.lab=2)
lines(uv$psi$smoothed.line, lwd=4)
abline(coef=c(0,1), lty=3)

# plot the derivative of correspondence curve on the scale of percentage
```

```

plot(uv$dpsi$t, uv$dpsi$value, xlab="t", ylab="psi'", xlim=c(0, max(uv$dpsi$t)),
ylim=c(0, max(uv$dpsi$value)), cex.lab=2)
lines(uv$dpsi$smoothed.line, lwd=4)
abline(h=1, lty=3)

# plot correspondence curve on the scale of the number of observations
plot(uv$psi.n$t, uv$psi.n$value, xlab="t", ylab="psi", xlim=c(0, max(uv$psi.n$t)),
ylim=c(0, max(uv$psi.n$value)), cex.lab=2)
lines(uv$psi.n$smoothed.line, lwd=4)
abline(coef=c(0,1), lty=3)

# plot the derivative of correspondence curve on the scale of the number
# of observations
plot(uv$dpsi.n$t, uv$dpsi.n$value, xlab="t", ylab="psi'", xlim=c(0, max(uv$dpsi.n$t)),
ylim=c(0, max(uv$dpsi.n$value)), cex.lab=2)
lines(uv$dpsi.n$smoothed.line, lwd=4)
abline(h=1, lty=3)

# If the rank lists consist of a large number of ties at the bottom
# (e.g. the same low value is imputed to the list for the observations
# that appear on only one list), it may be desirable to plot only
# observations before hitting the ties. Then it can be plotted using the
# following
plot(uv$psi$t[1:uv$psi$jump.point], uv$psi$value[1:uv$psi$jump.point], xlab="t",
ylab="psi", xlim=c(0, max(uv$psi$t[1:uv$psi$jump.point])),
ylim=c(0, max(uv$psi$value[1:uv$psi$jump.point])), cex.lab=2)
lines(uv$psi$smoothed.line, lwd=4)
abline(coef=c(0,1), lty=3)

```

salmon

Salmon data

Description

This data is from Simonoff (1990, p 161). It concerns the size of the annual spawning stock and its production of new catchable-sized fish for 1940 through 1967 for the Skeena river sockeye salmon stock (in thousands of fish). It has three columns, year, spawners and recruits. It was speculated to consist of two different populations.

Usage

```
data(salmon)
```

Format

A data frame with 28 observations on the following 3 variables.

year a numeric vector of the year

spawners a numeric vector of the annual spawning stock

recruits a numeric vector of the production of new catchable-sized fish

Source

Data is salmon.dat in Simonoff (1990). It can be downloaded from the book's website.

References

Simonoff, J. S. (1990), Smoothing Methods in Statistics, New York: Springer-Verlag.

Examples

```
data(salmon)
plot(rank(salmon$spawners), rank(salmon$recruits))
```

select.IDR	<i>Select observations according to IDR</i>
------------	---

Description

Select observations that exceeding a given IDR level

Usage

```
select.IDR(x, IDR.x, IDR.level)
```

Arguments

x	a n by m numeric matrix, where m= num of replicates, n=num of observations. Numerical values representing the significance of the observations, where larger values represent higher significance, for example, -log(p-value). Currently, m=2.
IDR.x	Irreproducible discovery rate for each entry of x. It is computed from est.IDR().
IDR.level	IDR cutoff, a numerical value between [0,1].

Value

x	Observations that are selected.
n	Number of observations that are selected.
IDR.level	IDR cutoff, a numerical value between [0,1].

Author(s)

Qunhua Li

References

Q. Li, J. B. Brown, H. Huang and P. J. Bickel. (2011) Measuring reproducibility of high-throughput experiments. Annals of Applied Statistics, Vol. 5, No. 3, 1752-1779.

See Also[est.IDR](#)**Examples**

```
data("simu.idr")
x <- cbind(-simu.idr$x, -simu.idr$y)

mu <- 2.6
sigma <- 1.3
rho <- 0.8
p <- 0.7

idr.out <- est.IDR(x, mu, sigma, rho, p, eps=0.001, max.ite=20)
# select observations exceeding IDR threshold=0.01
IDR.level <- 0.01
x.selected <- select.IDR(x, idr.out$IDR, IDR.level)
```

simu.idr

*Simulated data***Description**

This is a simulated dataset for testing the program. Data is first simulated from the copula mixture model with latent structure of $0.65 N(\mu, \mu, \sigma, \sigma, \rho) + 0.95 N(0, 0, 1, 1, 0)$, where $\mu=2.5$, $\sigma=1$, $\rho=0.84$. The observations in the dataset are then generated by taking the p-values from a z-test $H_0: \mu=0$.

Usage

```
data(simu.idr)
```

Format

A data frame with 1000 observations on the following 3 variables.

x a numeric vector, representing p-values on replicate 1

y a numeric vector, representing p-values on replicate 2

labels a binary vector, where 1 represents the reproducible component and 0 represents the irreproducible component.

References

Q. Li, J. B. Brown, H. Huang and P. J. Bickel. (2011) Measuring reproducibility of high-throughput experiments. *Annals of Applied Statistics*, Vol. 5, No. 3, 1752-1779.

Examples

```
data(simu.idr)
plot(rank(simu.idr$x), rank(simu.idr$y))
```


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