# Package 'QRank'

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Type Package
Title A Novel Quantile Regression Approach for eQTL Discovery
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<b>Description</b> A Quantile Rank-score based test for the identification of expression quantitative trait loci.
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QRank-package A Novel Quantile Regression Tool for eQTL Discovery

# Description

A Quantile Rank-score (QRank) based test for the identification of expression quantitative trait loci (eQTLs).

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#### **Details**

We use a Quantile Rank-score (QRank) based test to identify the expression quantitative trait loci (eQTLs) that are associated with the conditional quantile function of gene expressions.

#### Author(s)

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#### References

Xiaoyu Song, Gen Li, Zhenwei Zhou, Xianling Wang, Iuliana Ionita-Laza and Ying Wei (2016). QRank: A Novel Quantile Regression Tool for eQTL Discovery. *Under revision for Bioinformatics*.

## **Examples**

heter.QRank

Heterogeneity index for "QRank"

## **Description**

Calculate the heterogeneity index of quantile regression coefficients at multiple quantile levels. It measures the variation of the quantile coefficients across quantile levels.

```
heterogeneity = log(sd(\beta)/abs(mean(\beta)))
```

where  $\beta$  is the vector of quantile regression coefficients at multiple quantile levels.

#### Usage

```
heter.QRank(object, newtaus=NULL)
```

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#### **Arguments**

object Object returned from "QRank"

newtaus a vector of quantile levels based on which heterogeneity index are calculated.

Default is NULL, in which case the quantile levels inherited from "QRank" will

be used.

#### Value

```
heterogeneity index one hetergeneity index
```

## See Also

QRank

## **Examples**

```
# continuted from "QRank"

taus=c( 0.25, 0.5, 0.75)
q = QRank(gene=y, snp=x, cov=z, tau=taus)
heter.QRank(q) # default uses taus inherited from "QRank"
# - output
#Heterogeneity index:
#[1] 2.474184
heter.QRank(q,newtaus = 1:9/10) # calculate based on new taus values
# - output
#Heterogeneity index:
#[1] 2.69242
```

print.QRank

Print a QRank object

## **Description**

Print the object of QRank

## Usage

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

#### **Arguments**

x Object returned from QRank.

... Optional arguments

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#### See Also

QRank

print.QRank.heter

Print a QRank.heter object

## **Description**

Print the object of heter.QRank

## Usage

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

## Arguments

x Object returned from heter.QRank.

... Optional arguments

## See Also

heter.QRank

QRank

A new Quantile Rank-score (QRank) based test for the eQTL identification.

# Description

A function to obtain the p-value on the association between a gene expression and a genetic variant based on quantile rank-score test.

## Usage

```
QRank(gene, snp, cov = NULL, tau)
```

## Arguments

gene	a gene expression level from a selected gene. No parametric assumption is
	needed for underlying distribution.

snp a selected SNP.

cov a vector or matrix of covariates. Default is NULL.

tau the quantile levels to be estimated. Tau can be a single value or a vector of

quantile levels.

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## **Details**

This function conducts Quantile Rank-score (QRank) based test for the continuous traits. It can be used to identify expression quantitative trait loci (eQTLs) that are associated with the conditional quantile functions of gene expression.

#### Value

composite.pvalue

a single p-value for across all quantile levels under consideration, testing H0: No genetic association at the selected quantile levels.

quantile.specific.pvalue

p-values of each quantile level, testing  $H_0$ : The genetic variant and gene expression are not associated at this quantile level.

#### Author(s)

Xiaoyu Song

#### References

Xiaoyu Song, Gen Li, Zhenwei Zhou, Xianling Wang, Iuliana Ionita-Laza and Ying Wei (2016). QRank: A Novel Quantile Regression Tool for eQTL Discovery. *Under revision for Bioinformatics*.

## **Examples**

```
set.seed(123)
                                                                                     #
n=300
                                                                                     #
x=rbinom(n, 2, 0.2)
y=rnorm(n, mean=0, sd=1)
z=cbind(rbinom(n, 1, 0.3), rnorm(n, mean=2, sd=2))
taus=c( 0.25, 0.5, 0.75)
# - run the proposed QRank approach
                                                                                     #
QRank(gene=y, snp=x, cov=z, tau=taus)
                                                                                     #
# - output
                                                                                      #
#Composite.pvalue:
#[1] 0.2241873
#Quantile.specific.pvalue:
                                                                                     #
   0.25 0.5
                                                                                     #
#0.5452044 0.1821452 0.5938421
```

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