

Package ‘gQTLstats’

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Title gQTLstats: computationally efficient analysis for eQTL and allied studies

Version 1.2.0

Author VJ Carey <stvjc@channing.harvard.edu>

Description computationally efficient analysis of eQTL, mQTL, dsQTL, etc.

Suggests geuvPack, geuvStore, Rsamtools, knitr, rmarkdown, ggbio, BiocStyle, Homo.sapiens

Depends R (>= 3.1.0)

Imports methods, snpStats, BiocGenerics, S4Vectors, IRanges, GenomeInfoDb, GenomicRanges, SummarizedExperiment, VariantAnnotation, Biobase, BatchJobs, gQTLBase, limma, gam, dplyr, AnnotationDbi, GenomicFeatures, ggplot2, reshape2, doParallel, foreach, ffbase

Maintainer VJ Carey <stvjc@channing.harvard.edu>

License Artistic-2.0

LazyLoad yes

VignetteBuilder knitr

BiocViews SNP, GenomeAnnotation, Genetics

NeedsCompilation no

R topics documented:

gQTLstats-package	2
cisAssoc	3
clipPCs	5
directPlot	6
enumerateByFDR	7
eqBox2	8
FDRsupp-class	8
filtFDR	9
hmm878	10
manhWngr	12

queryVCF	13
senstab	14
setFDRfunc	15
storeToStats	16
txsPlot	18

Index	19
--------------	-----------

gQTLstats-package	<i>gQTLstats: computationally efficient analysis for eQTL and allied studies</i>
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Description

computationally efficient analysis of eQTL, mQTL, dsQTL, etc.

Details

The DESCRIPTION file:

```

Package:      gQTLstats
Title:        gQTLstats: computationally efficient analysis for eQTL and allied studies
Version:      1.2.0
Author:       VJ Carey <stvjc@channing.harvard.edu>
Description:  computationally efficient analysis of eQTL, mQTL, dsQTL, etc.
Suggests:    geuvPack, geuvStore, Rsamtools, knitr, rmarkdown, ggbio, BiocStyle, Homo.sapiens
Depends:      R (>= 3.1.0)
Imports:      methods, snpStats, BiocGenerics, S4Vectors, IRanges, GenomeInfoDb, GenomicRanges, SummarizedExp
Maintainer:   VJ Carey <stvjc@channing.harvard.edu>
License:      Artistic-2.0
LazyLoad:    yes
VignetteBuilder: knitr
BiocViews:    SNP, GenomeAnnotation, Genetics

```

Index of help topics:

```

FDRsupp-class      Class '"FDRsupp"'
cisAssoc           test for variant-expression associations in
                  cis, using VCF
clipPCs           transformations of expression data in smlSet
                  instances
directPlot        visualize relationship between empirical and
                  modeled FDR based on analysis of a gQTL store
enumerateByFDR    filter a ciseStore instance using an FDR
                  threshold
eqBox2            visualization of expression or other assay
                  measure against genotypes extracted from VCF

```

filtFDR	illustration of FDRsupp class
gQTLstats-package	gQTLstats: computationally efficient analysis for eQTL and allied studies
hmm878	labeled GRanges with ChromHMM chromatin states for GM12878
manhWngr	manhattan plot with named GRanges
queryVCF	obtain SnpMatrix from VCF genotypes
senstab	create a plottable table for eQTL sensitivity analysis visualization
setFDRfunc	estimate and store function relating association scores to approximate plug-in FDR
storeToQuantiles	extract a vector from store results as ff (out of memory reference); support statistical reductions
txsPlot	visualize transformed FDR against transformed association statistics

This package addresses the management of map-reduce like computations for cis-association tests between DNA variants and genomic features like gene expression measurements. It makes essential use of data structures defined in package gQTLBase.

A number of experimental functions are present in the current version of the package: prep.cisAssocNB (assembles information to assess negative binomial regression in cis association testing), storeToMaxAssocBySNP (progress towards SNP-specific FDR), table_sensobj_thresh (reporting on sensitivity analysis).

Author(s)

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cisAssoc *test for variant-expression associations in cis, using VCF*

Description

test for variant-expression associations in cis, using VCF and RangedSummarizedExperiment representations

Usage

```
cisAssoc(summex, vcf.tf, rhs = ~1, nperm = 3, cisradius =
  50000, genome = "hg19", assayind = 1, lbmaf = 1e-06,
  lbgtf = 1e-06, dropUnivHet = TRUE, infoFields =
  c("LDAF", "SVTYPE"), simpleSNV = TRUE)
cisCount(summex, vcf.tf, rhs = ~1, cisradius =
  50000, genome = "hg19", assayind = 1, lbmaf = 1e-06,
  lbgtf = 1e-06, dropUnivHet = TRUE, infoFields =
  c("LDAF", "SVTYPE"), simpleSNV = TRUE)
```

Arguments

summex	a RangedSummarizedExperiment object
vcf.tf	instance of TabixFile , referring to a tabix-indexed, bgzipped VCF file
rhs	formula ‘right hand side’ for adjustments to be made as snp.rhs.tests is run on each expression vector
nperm	number of permutations to be used for plug-in FDR computation
cisradius	distance in bp around each gene body to be searched for SNP association
genome	tag suitable for use in GenomeInfoDb structures
assayind	index of assays(summex) to use for expression data retrieval
lbmaf	lower bound on MAF of SNP to retain for analysis, computed using col.summary
lbgtf	lower bound on genotype frequency of SNP to retain for analysis
dropUnivHet	logical, if TRUE, will check for columns of SnpMatrix instance that possess no values other than "NA" and "A/B". See http://www.biostars.org/p/117155/#117270
infoFields	character – VCF fields to retain in vcfInfo() part of query
simpleSNV	logical – will use simple computation of isSNV to filter variants for analysis to SNV

Details

[snp.rhs.tests](#) is the workhorse for statistical modeling. VCF content is transformed to the byte-code (which allows for uncertain imputation) and used in fast testing.

Value

cisAssoc: a [GRanges-class](#) instance with mcols including chisq, permScore...
 cisCount: enumerate locations in VCF that would be tested

Note

seqlevelsStyle for summex and vcf.tf content must agree

Author(s)

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Examples

```
require(GenomeInfoDb)
require(geuvPack)
require(Rsamtools)
data(geuFPKM)
lgeu = geuFPKM[ which(seqnames(geuFPKM)=="chr20"), ]
seqlevelsStyle(lgeu) = "NCBI"
tf20 = TabixFile(system.file("vcf/c20exch.vcf.gz", package="GGtools"))
if (require(VariantAnnotation)) scanVcfHeader(tf20)
```

```

lgeue = clipPCs(lgeue[,which(lgeue$popcode=="CEU")], 1:2)
set.seed(1234)
litc = cisAssoc(lgeue[c(162,201),], tf20, nperm=2, lbmaf=.05, cisradius=50000)
summary(litc$chisq)
# \dontrun{
litc$pifdr = gQTLstats:::pifdr(litc$chisq, c(litc$permScore_1, litc$permScore_2))
litc[which(litc$pifdr < .01)]
# }

```

clipPCs

transformations of expression data in smlSet instances

Description

transformations of expression data in `smlSet` instances or assay data in `RangedSummarizedExperiment`

Usage

```
clipPCs(x, inds2drop, center = TRUE)
```

```
regressOut(x, rhs, ...)
```

Arguments

<code>x</code>	a <code>RangedSummarizedExperiment</code> object
<code>inds2drop</code>	Vector of PCs to be eliminated by setting the associated diagonal elements in the SVD to zero before recomposing the matrix of expression values. If the value 0 is present in <code>inds2drop</code> , the <code>smlSet</code> is returned unchanged, with a message.
<code>center</code>	logical, passed to <code>prcomp</code>
<code>rhs</code>	formula fragment (no dependent variable) used to form residuals in a reexpression of the expression matrix; variable bindings found in <code>pData</code> of an <code>ExpressionSet</code> or <code>colData</code> of a <code>RangedSummarizedExperiment</code>
<code>...</code>	arguments passed to <code>lmFit</code>

Details

`clipPCs` is an operation on the $n \times p$ transposed matrix X of expression data. The singular value decomposition $X = UDV^t$ is formed, the diagonal elements of D corresponding to `inds2drop` are set to zero yielding the diagonal matrix E , and then $Y = UEV^t$ is computed and transposed to replace the expression data.

`regressOut` obtains residuals after genewise regression of expression on the design matrix specified by the `rhs`; `lmFit` is used to compute coefficients, linear predictions and residuals.

Value

a `RangedSummarizedExperiment` object

Author(s)

VJ Carey <stvjc@channing.harvard.edu>

References

The use of PCA-based adjustments to remove mass extraneous effects from expression matrices has been criticized in work of Oliver Stegle and Jeffrey Leek, who offer Bayesian PEER and SVA respectively as alternative solutions.

Examples

```
if(require(geuvPack)){
  data(geuFPKM)
  cg = clipPCs(geuFPKM, 1:10)
  ro = regressOut(cg, ~popcode)
  ro
}
```

directPlot	<i>visualize relationship between empirical and modeled FDR based on analysis of a gQTL store</i>
------------	---

Description

visualize relationship between empirical and modeled FDR based on analysis of a gQTL store

Usage

```
directPlot(FDRsupp)
```

Arguments

FDRsupp instance of [FDRsupp-class](#)

Details

This plot is used to show the degree of fit between a smooth model relating modeled FDR to empirical FDR, and the empirical FDR themselves. It should be used in conjunction with [txsPlot](#).

It is possible for an implausible squiggly model to yield perfect agreement for all empirical FDR estimates. See the example.

Examples

```
data(filtFDR)
directPlot(filtFDR)
```

enumerateByFDR *filter a ciseStore instance using an FDR threshold*

Description

filter a ciseStore instance using an FDR threshold

Usage

```
enumerateByFDR(store, fdrsupp, threshold = 0.05, filter=force,
               ids=NULL, trimToUnit=TRUE)
```

Arguments

store	instance of ciseStore-class
fdrsupp	instance of FDRsupp-class
threshold	upper bound on FDR to be included
filter	The FDR can be computed for any association score. To return only records satisfying a given filter, supply the filter function here. It may be desirable to carry a filter function from the storeToFDR stage, and this may be considered in future versions.
ids	if NULL, process all results in store, otherwise limit attention to jobs with id values in ids
trimToUnit	plug-in FDR estimates can sometimes lie outside [0,1] owing to sparsity or defects of extrapolation; if this parameter is TRUE, estimated FDR values outside [0,1] are moved to the nearest boundary

Details

uses [storeApply](#), which will use BiocParallel infrastructure when available

Value

A GRanges instance with store contents to which estFDR is appended for each range. The estFDR quantity is predicted using the GAM model held in the FDRsupp instance.

Examples

```
require(geuvStore)
require(gQTLBase)
st = makeGeuvStore()
data(filtFDR)
filtEnum = enumerateByFDR( st, filtFDR,
  filter=function(x)x[which(x$mindist <= 500000 & x$MAF >= 0.05)] )
names(metadata(filtEnum))
filtEnum[order(filtEnum$chisq, decreasing=TRUE)[1:2]]
```

eqBox2	<i>visualization of expression or other assay measure against genotypes extracted from VCF</i>
--------	--

Description

visualization of expression or other assay measure against genotypes extracted from VCF

Usage

```
eqBox2(gene, se, tf, snpgr, genome = "hg19", ...)
eqDesc2(gene, se, tf, snpgr, genome = "hg19")
```

Arguments

gene	an element of rownames(se) from which a vector of assay values will be created
se	a RangedSummarizedExperiment object
tf	instance of class TabixFile-class , defining paths to a tabix-indexed VCF and index file
snpgr	instance of GRanges-class identifying the SNP to be visualized
genome	tag identifying reference genome
...	extra arguments passed to boxplot

Examples

```
require(Rsamtools)
require(SummarizedExperiment)
mygr = GRanges("1", IRanges(54683925, width=1))
gene = "ENSG00000231581.1"
library(geuvPack)
data(geuFPKM)
#tf = gtpath(1)
tf = TabixFile(system.file("vcf/small_1.vcf.gz", package="gQTLstats"))
eqBox2(gene, se=geuFPKM, tf, mygr )
eqDesc2(gene, se=geuFPKM, tf, mygr )
```

FDRsupp-class	<i>Class "FDRsupp"</i>
---------------	------------------------

Description

Support for FDR computations with ciseStore instances

Objects from the Class

Objects can be created by calls of the form `new("FDRsupp", ...)`.

Slots

tab: Object of class "data.frame" a table with association scores and plug-in FDR estimates evaluated on selected score values

FDRfunc: Object of class "function" a function of one argument with input association score and output the corresponding FDR estimate

FDRmodel: Object of class "gam" that was fit to elements of tab

filterUsed: Object of class "function" a copy of the function used for filtering the store to create the FDRfunc element.

sessinfo: `sessionInfo()` value at time of construction

theCall: instance of class "call" showing call leading to construction

Methods

getFDRfunc signature(`x = "FDRsupp"`): extract the FDR approximating function, a function of one (vector) argument assumed to represent association scores, evaluating to the plug-in FDR estimates corresponding to these scores

getTab signature(`x = "FDRsupp"`): extract the table of association scores and empirical FDR estimates

Note

Typically the `FDRfunc` function is constructed using a smooth model relating the estimated FDR to association scores.

Examples

```
showClass("FDRsupp")
```

`filtFDR`

illustration of FDRsupp class

Description

illustration of FDRsupp class

Usage

```
data("filtFDR")
```

Format

A FDRsupp object.

Details

filtFDR was constructed on geuvStore contents, filtering to MAF at least five percent and radius at most 500kbp. rawFDR uses the entire geuvStore contents, with 1Mbp radius and 1 percent MAF lower bound

Examples

```
data(filtFDR)
filtFDR
```

hmm878

labeled GRanges with ChromHMM chromatin states for GM12878

Description

labeled GRanges with ChromHMM chromatin states for GM12878

Usage

```
data(hmm878)
```

Format

The format is:

```
Formal class 'GRanges' [package "GenomicRanges"] with 6 slots
..@ seqnames :Formal class 'Rle' [package "IRanges"] with 4 slots
.. ..@ values : Factor w/ 23 levels "chr1","chr2",...: 1 2 3 4 5 6 7 8 9 10 ...
.. ..@ lengths : int [1:23] 54467 46499 37617 25155 30071 34846 29420 24506 24123 27263 ...
.. ..@ elementMetadata: NULL
.. ..@ metadata : list()
..@ ranges :Formal class 'IRanges' [package "IRanges"] with 6 slots
.. ..@ start : int [1:571339] 10001 10601 11138 11738 11938 12138 14538 20338 22138 22938
...
.. ..@ width : int [1:571339] 600 537 600 200 200 2400 5800 1800 800 4000 ...
.. ..@ NAMES : NULL
.. ..@ elementType : chr "integer"
.. ..@ elementMetadata: NULL
.. ..@ metadata : list()
..@ strand :Formal class 'Rle' [package "IRanges"] with 4 slots
.. ..@ values : Factor w/ 3 levels "+","-","*": 3
.. ..@ lengths : int 571339
.. ..@ elementMetadata: NULL
.. ..@ metadata : list()
..@ elementMetadata:Formal class 'DataFrame' [package "IRanges"] with 6 slots
.. ..@ rownames : NULL
.. ..@ nrows : int 571339
.. ..@ listData :List of 4
```

```

.. .. ..$ name : chr [1:571339] "15_Repetitive/CNV" "13_Heterochrom/lo" "8_Insulator" "11_Weak_Txn"
...
.. .. ..$ score : num [1:571339] 0 0 0 0 0 0 0 0 0 ...
.. .. ..$ itemRgb: chr [1:571339] "#F5F5F5" "#F5F5F5" "#0ABEFE" "#99FF66" ...
.. .. ..$ thick :Formal class 'IRanges' [package "IRanges"] with 6 slots
.. .. .. .. ..@ start : int [1:571339] 10001 10601 11138 11738 11938 12138 14538 20338 22138
22938 ...
.. .. .. .. ..@ width : int [1:571339] 600 537 600 200 200 2400 5800 1800 800 4000 ...
.. .. .. .. ..@ NAMES : NULL
.. .. .. .. ..@ elementType : chr "integer"
.. .. .. .. ..@ elementMetadata: NULL
.. .. .. .. ..@ metadata : list()
.. .. ..@ elementType : chr "ANY"
.. .. ..@ elementMetadata: NULL
.. .. ..@ metadata : list()
..@ seqinfo :Formal class 'Seqinfo' [package "GenomicRanges"] with 4 slots
.. .. ..@ seqnames : chr [1:23] "chr1" "chr2" "chr3" "chr4" ...
.. .. ..@ seqlengths : int [1:23] 249250621 243199373 198022430 191154276 180915260 171115067
159138663 146364022 141213431 135534747 ...
.. .. ..@ is_circular: logi [1:23] FALSE FALSE FALSE FALSE FALSE FALSE ...
.. .. ..@ genome : chr [1:23] "hg19" "hg19" "hg19" "hg19" ...
..@ metadata :List of 1
.. ..$ url: chr "http://genome.ucsc.edu/cgi-bin/hgFileUi?g=wgEncodeBroadHmm&db=hg19"

```

Details

acquired using rtracklayer import from the bed file given at metadata(hmm878)[["url"]]

Source

see details

References

Ernst J, Kellis M. Discovery and characterization of chromatin states for systematic annotation of the human genome. *Nat Biotechnol.* 2010 Aug;28(8):817-25.

Ernst J, Kheradpour P, Mikkelsen TS, Shores N, Ward LD, Epstein CB, Zhang X, Wang L, Issner R, Coyne M et al. Mapping and analysis of chromatin state dynamics in nine human cell types. *Nature.* 2011 May 5;473(7345):43-9.

Examples

```

data(hmm878)
table(hmm878$name)

```

manhWngr	<i>manhattan plot with named GRanges</i>
----------	--

Description

manhattan plot with named GRanges

Usage

```
manhWngr(store, probeid = "ENSG00000183814.10", sym = "LIN9", fdrsupp, namedGR, slstyle = "NCBI", xlab
```

Arguments

store	instance of ciseStore-class
probeid	name of feature identifier to use for cis association
sym	symbol for feature identifier
fdrsupp	instance of FDRsupp-class
namedGR	GRanges instance with 'name' in mcols element
slstyle	seqlevelsStyle
xlab.in	x axis label
ylab.in	y axis label
applyFDRfilter	if TRUE, use the filter defined in the filterUsed element of the object supplied as fdrsupp on the output
...	additional arguments for plotting

Examples

```
require(geuvStore)
require(gQTLBase)
store = makeGeuvStore()
data(hmm878)
data(filtFDR)
manhWngr(store, fdrsupp=filtFDR, namedGR=hmm878)
```

queryVCF	<i>obtain SnpMatrix from VCF genotypes</i>
----------	--

Description

obtain SnpMatrix from VCF genotypes

Usage

```
queryVCF(gr, vcf.tf, samps, genome = "hg19", getSM = TRUE)
```

Arguments

gr	GRanges instance; SNPs lying within will be processed
vcf.tf	TabixFile instance pointing to VCF
samps	samples to be retained
genome	tag identifying build
getSM	logical; if FALSE, genotypeToSnpMatrix will not be run and only the output of readVcf is returned.

Value

a list of length two

readout	output of readVcf
sm	output of genotypeToSnpMatrix run on the read result

Examples

```
require(Rsamtools)
tf20 = TabixFile(system.file("vcf/c20exch.vcf.gz", package="GGtools"))
require(geuvPack)
data(geuFPKM)
lgeu = geuFPKM[ which(seqnames(geuFPKM)=="chr20"),
               which(geuFPKM$popcode=="CEU") ]
seqlevelsStyle(lgeu) = "NCBI"
rng = rowRanges(lgeu)[232] # CPNE1
myq = queryVCF( rng, tf20, samps=colnames(lgeu), genome="hg19" )
myq
```

senstab	<i>create a plottable table for eQTL sensitivity analysis visualization</i>
---------	---

Description

create a plottable table for eQTL sensitivity analysis visualization

Usage

```
senstab(x, filt = force)
## S3 method for class 'senstab'
plot(x, ...)
```

Arguments

x	a list generated by a process analogous to the sensitivity survey exhibited in the example below
filt	a function that operates on and returns a data.frame; typically will select rows based on values of fields 'MAF' and 'radius'
...	extra arguments passed to plot

Details

sensByProbe is a list structure; for information on this and other elements of sensitivity analysis workflow, see extensive non-executed code in example below

Value

an instance of the S3 class 'senstab', 'data.frame'

Examples

```
## Not run:
#
# illustration of sensitivity analysis using BatchJobs
#
# assume the following content in 'parms.R' (uncommented)
# MAFS = c(.03, .04, .05, .075, .10, .125, .15)
# dists = c(5000, 7500, 10000, 15000, 20000,
# 25000, 50000, 100000, 250000, 500000, 750000, 1000000)
# parms = expand.grid(MAFS, dists)
library(BatchJobs) # for bigStore manip
library(gQTLstats)

# could use multilevel parallelism here
# because it is a somewhat large, fragile job, BatchJobs
# is a relevant tool for iteration. but storeToFDRByProbe is
# already using bplapply. so register 3 cores for it
```

```

# and specify 15 cpu for BatchJobs in .BatchJobs.R

sens1 = makeRegistry("sens1", file.dir="sens1",
  packages=c("gQTLstats", "dplyr"),
  src.files="parms.R") # note parms.R

sens4One = function(z) {
  load("../bigStore.rda") # get a ciseStore instance
  ans = storeToFDRByProbe(bigStore, xprobs=seq(.01,.99,.01), # xprobs
    # needs to be chosen with care
  filter=function(x) x[which(x$MAF >= parms[z,1] &
    x$mindist <= parms[z,2])])
  ans = setFDRfunc(ans, span=.35) # span can be important
  list(fdrsups=ans, parms=parms[z,])
}

batchMap(sens1, sens4One, 1:nrow(parms))
submitJobs(sens1)

# now loadResult(sens1) or the equivalent can be the input to sensstab()
# as in the example to continue here:

## End(Not run)
library(gQTLstats)
data(sensByProbe)
ptab = t(sapply(sensByProbe, function(x)as.numeric(x[[2]])))
unique(ptab[,1]) # MAFs used
unique(ptab[,2]) # radii used
# here we filter away some extreme values of the design space
tab = sensstab(sensByProbe, filt=function(x) {
  x[ x$radius > 10000 & x$ radius < 500000 & x$MAF > .03, ]
} )
plot(tab)

```

setFDRfunc	<i>estimate and store function relating association scores to approximate plug-in FDR</i>
------------	---

Description

estimate and store function relating association scores to approximate plug-in FDR

Usage

```
setFDRfunc(FDRsupp, fudge = 1e-06, zthresh = 30, ...)
```

Arguments

FDRsupp instance of [FDRsupp-class](#)

fudge	if FDR is zero, a log or logistic transform will fail; we add the small positive number fudge to avoid this
zthresh	for association scores greater than this value, a hard value of FDR 0 is assigned
...	arguments passed to <code>lo</code> for the smooth model relating association score to FDR at selected quantiles of the association score distribution

Value

returns an updated `FDRsupp-class` instance

Examples

```
data(filtFDR)
filtFDR2 = setFDRfunc(filtFDR)
```

storeToStats	<i>extract a vector from store results as ff (out of memory reference); support statistical reductions</i>
--------------	--

Description

extract a vector from store results as ff (out of memory reference); support statistical reductions

Usage

```
storeToQuantiles(store, field,
  probs=c(seq(0, .999, .001), 1-(c(1e-4,1e-6,1e-6,1e-7))),
  ids = NULL, ..., checkField = FALSE, filter=force)
storeToHist(store, getter = function(x)
  as.numeric(S4Vectors::as.matrix(mcols(x)[,
  grep("permScore", names(mcols(x)))])), breaks, ids =
  NULL, filter = force)
storeToFDR(store, xprobs = c(seq(0, 0.999, 0.001), 1 - (c(1e-04,
  1e-06, 1e-06, 1e-07))), xfield = "chisq", getter =
  function(x) as.numeric(S4Vectors::as.matrix(mcols(x)[,
  grep("permScore", names(mcols(x)))])), filter = force)
```

Arguments

store	instance of <code>ciseStore-class</code>
field	character tag, length one, must be name of a numeric field in the result set (typically something like 'chisq' in the GRanges generated by <code>cisAssoc</code>)
xfield	as field, for FDR computation, see Details.
ids	job ids to be used; if NULL, process all jobs
breaks	boundaries of histogram bins

...	supplied to makeRegistry for a temporary registry: typically will be a vector of package names if additional packages are needed to process results
checkField	if TRUE steps will be taken to verify that the tag to which 'field' evaluates is present in result in the first job
probs	numeric vector of probabilities with values in [0,1]. See quantile.ff .
xprobs	percentiles of the empirical distribution of the association statistic at which FDR estimates are recorded.
getter	function of a single argument that extracts a numeric vector of association scores obtained under permutation
x	instance of FDRsupp
filter	function accepting and returning GRanges instance, executed when cisAssoc result is loaded to modify that result, defaults to no-op

Details

uses current BatchJobs configuration to parallelize extraction; reduceResults could be used for a sequential solution

Value

storeToQuantiles and storeToHist return objects analogous to those returned by stats::quantile and graphics::hist.

However, it should be noted that storeToQuantiles will use the [quantile.ff](#) of ffbase. For vectors of modest length, this can disagree with results of base::quantile by a few percent.

storeToFDR and storeToFDRByProbe return an instance of FDRsupp class

Note

uses ffbase:::c.ff explicitly to concatenate outputs; there is no guarantee of order among elements

Examples

```
stopifnot(require(geuvStore))
require(BatchJobs)
require(gQTLBase)
store = makeGeuvStore()
library(doParallel)
if (.Platform$OS.type == "windows") {
  registerDoSEQ()
} else registerDoParallel(cores=max(c(detectCores()-1,1)))
smchisq = storeToFf( store, "chisq", ids=store@validJobs[1:3])
smchisq
qs = storeToQuantiles( store, "chisq", ids = store@validJobs[1:5],
  probs=seq(.1,.9,.1) )
qs
hh = storeToHist( store, ids = store@validJobs[1:5], breaks=
  c(0,qs,1e9) )
hh$counts
```

```
fd = storeToFDR( store, xprobs=c(seq(.05,.95,.05),.99,.999) )
tail(getTab(fd),4)
sss = storeToFDRByProbe( store , xprobs=c(seq(.05,.95,.05),.99) )
tail(getTab(sss),4)
```

txsPlot*visualize transformed FDR against transformed association statistics*

Description

visualize transformed FDR against transformed association statistics

Usage

```
txsPlot(FDRsupp)
```

Arguments

FDRsupp an instance of [FDRsupp-class](#)

Examples

```
data(filtFDR)
txsPlot(filtFDR)
```

Index

- *Topic **classes**
 - FDRsupp-class, 8
- *Topic **datasets**
 - filtFDR, 9
 - hmm878, 10
- *Topic **graphics**
 - directPlot, 6
 - eqBox2, 8
 - txsPlot, 18
- *Topic **models**
 - cisAssoc, 3
 - clipPCs, 5
 - enumerateByFDR, 7
 - manhWngr, 12
 - queryVCF, 13
 - senstab, 14
 - setFDRfunc, 15
 - storeToStats, 16
- *Topic **package**
 - gQTLstats-package, 2
- cisAssoc, 3
- cisCount (cisAssoc), 3
- clipPCs, 5
- clipPCs, RangedSummarizedExperiment, numeric, logical-method (clipPCs), 5
- clipPCs, RangedSummarizedExperiment, numeric, missing-method (clipPCs), 5
- clipPCs, SummarizedExperiment, numeric, logical-method (clipPCs), 5
- clipPCs, SummarizedExperiment, numeric, missing-method (clipPCs), 5
- col.summary, 4
- directPlot, 6
- enumerateByFDR, 7
- eqBox2, 8
- eqDesc2 (eqBox2), 8
- FDRsupp-class, 8
- filtFDR, 9
- genotypeToSnpMatrix, 13
- getFDRfunc (FDRsupp-class), 8
- getFDRfunc, FDRsupp-method (FDRsupp-class), 8
- getTab (FDRsupp-class), 8
- getTab, FDRsupp-method (FDRsupp-class), 8
- gQTLstats (gQTLstats-package), 2
- gQTLstats-package, 2
- GRanges, 12
- hmm878, 10
- isSNV, 4
- lmFit, 5
- lo, 16
- manhWngr, 12
- plot (senstab), 14
- prcomp, 5
- prep.cisAssocNB (gQTLstats-package), 2
- quantile.ff, 17
- queryVCF, 13
- RangedSummarizedExperiment, 4, 5, 8
- rawFDR (filtFDR), 9
- regressOut (clipPCs), 5
- setByProbe (senstab), 14
- senstab, 14
- setFDRfunc, 15
- snp.rhs.tests, 4
- storeApply, 7
- storeToFDR (storeToStats), 16
- storeToFDRByProbe (storeToStats), 16
- storeToHist (storeToStats), 16
- storeToMaxAssocBySNP (gQTLstats-package), 2

storeToQuantiles (storeToStats), [16](#)
storeToStats, [16](#)

TabixFile, [4](#)

table_sensobj_thresh
(gQTLstats-package), [2](#)

txsPlot, [6](#), [18](#)