Package ‘gQTLBase’

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Title  gQTLBase: infrastructure for eQTL, mQTL and similar studies
Version  1.6.0
Author  VJ Carey <stvjc@channing.harvard.edu>
Description  Infrastructure for eQTL, mQTL and similar studies.
Suggests  geuvStore2, knitr, rmarkdown, BiocStyle, RUnit, GGtools,
          Homo sapiens, IRanges, erma, GenomeInfoDb, gwascat, geuvPack
Imports  GenomicRanges, methods, BatchJobs, BBmisc, S4Vectors,
          BiocGenerics, foreach, doParallel, bit, ff, rtracklayer,
          ffbase, GenomicFiles, SummarizedExperiment
Depends
Maintainer  VJ Carey <stvjc@channing.harvard.edu>
License  Artistic-2.0
LazyLoad  yes
VignetteBuilder  knitr
BiocViews  SNP, GenomeAnnotation, Genetics, DataImport,
          FunctionalGenomics
          mergeToLoci.R ufeatByTiling.R d.R
NeedsCompilation  no

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gQTLBase-package

**gQTLBase: infrastructure for eQTL, mQTL and similar studies**

**Description**

Infrastructure for eQTL, mQTL and similar studies.

**Details**

The DESCRIPTION file:

- **Package:** gQTLBase
- **Title:** gQTLBase: infrastructure for eQTL, mQTL and similar studies
- **Version:** 1.6.0
- **Author:** VJ Carey <stvjc@channing.harvard.edu>
- **Description:** Infrastructure for eQTL, mQTL and similar studies.
- **Suggests:** geuvStore2, knitr, markdown, BiocStyle, RUnit, GGtools, Homo.sapiens, IRanges, erma, GenomeInfoDb
- **Imports:** GenomicRanges, methods, BatchJobs, BBmisc, S4Vectors, BiocGenerics, foreach, doParallel, bit, ff, rtracklayer
- **Depends:**
- **Maintainer:** VJ Carey <stvjc@channing.harvard.edu>
- **License:** Artistic-2.0
- **LazyLoad:** yes
- **VignetteBuilder:** knitr
- **BiocViews:** SNP, GenomeAnnotation, Genetics, DataImport, FunctionalGenomics

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- **ufeatByTiling**
  - split featurenames of SummarizedExperiment according to tiling, or to achieve simple balance within seqnames

Purpose is to define infrastructure on a comprehensive archive of eQTL, mQTL, dsQTL, etc., association statistics.
Description

wrap a BatchJobs registry that manages results of a cis-eQTL search

Objects from the Class

Objects can be created by calls of the form `new("ciseStore", reg=reg, ...)`. All arguments must be named.

We can also use `ciseStore(reg, validJobs, addProbeMap = TRUE, addRangeMap = TRUE)` and the probemap and rangeMap slots will be populated appropriately. If validJobs is missing, the validJobs slot will be populated by `findDone(reg)`. This may be problematic for handcrafted extracts from archives.

Slots

- `reg`: Object of class "Registry" BatchJobs Registry instance
- `validJobs`: Object of class "integer" vector of valid job identifiers for the registry
- `probemap`: Object of class "data.frame" a map from expression probe identifiers to job identifiers where results for the probe are stored
- `rangeMap`: Object of class "GRanges" a map from ranges on chromosomes, to job identifiers, in `mcols()$jobid`

Methods

- `show`

Function `describeStore` uses `batchMapResults` and `reduceResults` to leverage a parallel environment to collect information on numbers of tests and features. Arguments are described in the associated man page.

Note

the construction of the maps occurs via `storeApply`, which will use `foreach`, so that registration of a parallel back end using, e.g., `registerDoParallel`, will determine the speed of construction

Any registry job results that do not inherit from GRanges are mapped to NULL and will not be present in ultimate maps.

describeStore

collect basic descriptive statistics on ciseStore instances

Description

collect basic descriptive statistics on ciseStore instances

Usage

describeStore(st, genetag = "probeid", snptag = "snp", ids = NULL,
resfilter = force, doChecks = TRUE, ...)
describeByFilts(st, filtlist, ...)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>st</td>
<td>instance of <strong>ciseStore-class</strong></td>
</tr>
<tr>
<td>genetag</td>
<td>string for field name for name of quantitatively assayed feature, defaults to &quot;probeid&quot;; for GTEx application &quot;gene&quot; is used</td>
</tr>
<tr>
<td>snptag</td>
<td>string for field name for name of genotype feature</td>
</tr>
<tr>
<td>ids</td>
<td>integerish vector of ids, can be left NULL to survey entire store</td>
</tr>
<tr>
<td>resfilter</td>
<td>function applied to job results prior to summarization, defaults to force()</td>
</tr>
<tr>
<td>filtlist</td>
<td>a list of functions suitable as resfilter arguments</td>
</tr>
<tr>
<td>doChecks</td>
<td>logical – if true, will collect information on match between number probes requested and number reported on, and two scans of VCF loci in cis to probes. See details.</td>
</tr>
<tr>
<td>...</td>
<td>used with describeByFilts, pass to storeApply</td>
</tr>
</tbody>
</table>

Details

uses parallel infrastructure of foreach on contents managed by st@reg
describeByFilts returns a matrix of descriptions with one row per filtlist element
storeDescription holds results of a describe task and includes information on noncongruence of features with cis tests and of results of two distinct scans of VCF: one with readGT on a single sample, the other with readVcf on all samples. If there are discrepancies between features given and tests returned, [storeDescription]@reqfail will give the job ids for these. If there are discrepancies between the numbers of loci retrieved on the two VCF scans, @locfail will tive the job ids for these. @reqfail events may be legitimate when a feature has no SNP in cis at the given radius. @locfail events usually indicate an I/O problem and the jobs should be resubmitted.
Value

list with elements ntests, ngene.uniq, nsnp.uniq

Examples

## Not run:
library(geuvStore2)
mm = makeGeuvStore2()
describeStore(mm, ids=1:10, resfilter=function(x) x[!x$mindist < 50000])
## End(Not run)

extractByProbes retrieve eqtlTest results from a ciseStore instance

Description

retrieve eqtlTest results from a ciseStore instance

Usage

extractByProbes(store, probeids, extractTag = "probeid")
extractByRanges(store, gr)
extractBySymbols(store, symbols, sym2probe, extractTag = "probeid")

Arguments

store instance of ciseStore-class
probeids, symbols vector character tokens
gr instance of GRanges-class
sym2probe named character vector of probeids with names given by corresponding symbols
extractTag character atom telling what field in the archived GRanges is regarded as the probe or gene identifier
... extra arguments to extractByProbes

Details

an index will be searched if created by the ciseStore constructor

Value

a GRanges instance

Author(s)

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

merge ChromImpute chromatin states, or GWAS hit indicators, to a GRanges

Description

merge ChromImpute chromatin states, or GWAS hit indicators, to a GRanges

Usage

mergeCIstates(gr, ermaset=NULL, epig, genome = "hg19", importFull=FALSE, useErma = TRUE, stateGR=NULL)
mergeGWhts(gr, gwcat, use=c("both", "addr", "name")[1],
grSnpField="SNP")

Arguments

- **gr**: a GRanges instance
- **ermaset**: an instance of `ErmaSet-class`, if NULL, supply a GRanges as stateGR, with fields states and statecols
- **gwcat**: an instance of `gwaswloc-class`, or any compliant GRanges instance – must have mcols field SNPS with snp identifier
- **epig**: the standardized epigenome name of the epigenome to use
- **genome**: a tag for genome build
- **importFull**: logical, set to TRUE to acquire entire content (for LNG.FET, 800K ranges), to avoid contention for connections in parallel applications
- **useErma**: logical – at the moment, must be TRUE; plan is to allow use of elements of AnnotationHub
- **use**: character string selecting approach for linking loci in gr to those in gwcat – if "both", coincidence in address or name are both checked and used; if "addr", only address is checked, if "name", only SNP name.
- **grSnpField**: character string naming the field in mcols(gr) with SNP id
- **stateGR**: a GRanges instance as imported from erma package or from AnnotationHub, with mcols field states denoting chromatin state and statecols the associated colors for rendering
Value

for mergeCIstates, a GRanges instance with additional fields in mcols: fullStates, states, and state-
cols, denoting respectively the full annotation of ChromImpute for the inferred state, an abbreviated
tag that collapses related states, and a color tag for rendering, that does not replicate the colors in the
ChromImpute bed files. The states field is a factor with levels c("Het", "DNase", "Enh", "Prom", "Quies", "ReprPC", "Tss", "Tx","ZNF/Rp")
for mergeGWhits, a single mcols field is added, isGwasHit, that is 1 for coincident hit and 0
otherwise. Eventually phenotype information will be collected and added.

Examples

```r
if (require(gwascat) && require(erma)) {
  # demonstrate Tx state for exon starts
  gm = resize(genemodel("ORMDL3"),1)
  es = makeErmaSet()
  g1 = mergeCIstates(gm, es, "LNG.FET")
  g1

  # set up for GWAS
  #
  require(GenomeInfoDb)
  data(ebicat37)
  genome(ebicat37) = "hg19"
  seqlevelsStyle(ebicat37) = "UCSC"
  g1 = c(g1, g1[1]) # add a known hit
  start(g1[length(g1)]) = 38062196
  mergeGWhits(g1, ebicat37)
}
```

Description

apply a function over job results in a ciseStore instance

Usage

`storeApply(store, f, n.chunks, ids=NULL, ..., verbose = FALSE, flatten1=TRUE)`

Arguments

- **store**: instance of `ciseStore-class`
- **f**: function on GRanges stored in ciseStore
- **n.chunks**: Number of chunks into which the jobs are to be broken; the
  series of chunks is handed to `foreach` to extract results and apply `f` to them.
  If missing, the value of `getDoParWorkers()` used.
- **ids**: defaults to NULL; if non-null, the jobs to be processed are limited to those
  identified in this vector.
... additional arguments to foreach
verbose if TRUE will allow progressbars and other messages to display
flatten1 if TRUE will execute `unlist(...)`, defaulted to FALSE in previous version

Details

The chunking of job identifiers will determine the degree of parallelization of application, and the form of the list that is returned. `flatten1` will eventually default to TRUE.

Value

A list whose structure depends on the chunking of job identifiers. See the examples.

Note

eqtlStore imports Bioconductor’s bpparam function, and this determines in real time the number of workers to be employed by `storeApply`.

See Also

`storeMapResults` will apply over the store using the batch jobs submission infrastructure and can target specific results via `ids`; `storeApply` uses `bplapply` over the entire store.

Examples

```r
if (require(geuvStore2)) {
  require(BatchJobs)
  store = makeGeuvStore2()
  storeApply(store, length)
  storeApply(store, length, ids=c(1:3, 603))
}
```

---

use batchMapResults infrastructure to process results in a ciseStore instance

Description

use batchMapResults infrastructure to process results in a ciseStore instance

Usage

```r
storeMapResults(store, reg2, fun, ...,
ids = NULL, part = NA_character_, more.args = list())
loadAndFilterResult(reg,
id, filter=force, part = NA_character_, missing.ok = FALSE)
```
storeMapResults

Arguments

store  an instance of ciseStore-class
reg    instance of BatchJobs Registry class
reg2   an empty instance of the Registry class (see makeRegistry)
fun    A function to map over results in store, with formals (job, res, ...).
filter a function that accepts and returns a GRanges instance, to be applied just after
        loading a result from the store
...    additional arguments to vectorize over (should be same length as length(findDone(store@reg)))
ids    ids of job results to be mapped; if missing, map all job results
id     a single job id
part   see batchMapResults
missing.ok see loadResult
more.args a list of other arguments to be passed to fun; default is empty list.

Value

integer vector with job ids. Main purpose is to prepare the registry for submitJobs.

Note

loadAndFilterResult is not intended to be exported and may be removed in future versions.

Author(s)

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Examples

## Not run:
if (require(geuvStore2)) {
  require(BatchJobs)
  store = makeGeuvStore2()
  fd = tempfile()
  tempreg = makeRegistry("tempSMR", file.dir=fd)
  storeMapResults( store, tempreg, func=function(job, res, ...) length(res) )
  showStatus(tempreg)
  submitJobs(tempreg, 1:2)
  loadResults(tempreg)
  unlink(fd)
}

## End(Not run)
storeToFf  

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

Description

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

Usage

storeToFf(store, field, ids = NULL, filter=force, ..., checkField = FALSE, ischar=FALSE)

Arguments

store  
instance of ciseStore-class

field  
character tag, length one. If name of a numeric field in the result set (typically something like 'chisq' in the GRanges generated by cisAssoc), ff is applied directly. Character variables are converted to factors before ff is applied.

ids  
job ids to be used; if NULL, process all jobs

filter  
function to be applied when GRanges is loaded from results store, should accept and return a GRanges instance

...  
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

ischar  
must be true for character vector to be handled properly as a factor, otherwise NA will be returned

Details

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

Value

a vector as ff reference

Note

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

Examples

```r
if (require(geuvStore2)) {
  require(BatchJobs)
  store = makeGeuvStore2()
  smchisq = storeToFf( store, "chisq", ids=store@validJobs[1:3])
  smchisq
}
```
**ufeatByTiling**

**ufeatByTiling**  
*split featurenames of SummarizedExperiment according to tiling, or to achieve simple balance within seqnames*

**Description**  
split featurenames of SummarizedExperiment according to tiling, without redundancies

**Usage**
```
ufeatByTiling(se, tiling, maxlen=20)
balancedFeatList(se, maxlen=20)
```

**Arguments**
- `se`: instance of SummarizedExperiment
- `tiling`: GRanges instance corresponding to a genomic tiling
- `maxlen`: numeric ... list elements longer than maxlen are chopped up to have this length, to foster load balancing

**Details**

ufeatByTiling uses findOverlaps, balancedFeatList uses split on seqnames and BBmisc::chunk

**Value**

a list with elements of names(rowRanges(se)) corresponding to the elements of the tiling

**Examples**
```r
## Not run:
library(geuvPack)
data(geuFPKM)
library(Homo.sapiens)
au = paste0("chr", 1:22)
tg_500k = tileGenome(seqinfo(TxDb(Homo.sapiens))[au,], tilewidth=500000, cut.last.tile.in.chrom=TRUE)
sn = ufeatByTiling(geuFPKM, tg_500k)
summary(sapply(sn,length))
sn2 = balancedFeatList(geuFPKM)
summary(sapply(sn2,length))
## End(Not run)
```
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