Package 'gwascat'

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Title representing and modeling data in the EMBL-EBI GWAS catalog
Version 2.2.1
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Description Represent and model data in the EMBL-EBI GWAS catalog.
Enhances SNPlocs. Hsapiens.dbSNP.20120608
Depends R (>= 3.0.0), Homo.sapiens
Imports methods, BiocGenerics, S4Vectors, IRanges, GenomeInfoDb, GenomicRanges, snpStats, Biostrings, Rsamtools, rtracklayer, gQTLstats, Gviz, VariantAnnotation, AnnotationHub, AnnotationDbi, GenomicFeatures, graph, ggbio, ggplot2
Suggests DO.db, DT, utils, knitr, RBGL, RUnit, GGtools
VignetteBuilder utils, knitr
Maintainer VJ Carey <stvjc@channing.harvard.edu></stvjc@channing.harvard.edu>
License Artistic-2.0
LazyData no
biocViews Genetics
NeedsCompilation no

${\sf R}$ topics documented:

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gwascat-package representing and modeling data in the NHGRI GWAS catalog

Description

representing and modeling data in the NHGRI GWAS catalog, using GRanges and allied infrastructure

Details

Package: gwascat Version: 1.7.3

Suggests:

Depends: R (>= 3.0.0), methods, IRanges, GenomicRanges

Imports:

License: Artistic-2.0

LazyLoad: yes

Index:

The GWAS catalog management has migrated to EMBL/EBI. Use data(ebicat38) for an image dated 3 August 2015. Use makeCurrentGwascat() to get a more recent image. Use data(ebicat37) for a GRCh37 (or hg19) liftOver result. Use data(ebicat37UCSC) for an image with hg19 as genome tag and UCSC seqnames.

The data objects

```
'g17SM' 'gg17N' 'gw6.rs_17' 'low17' 'rules_6.0_1kg_17' 'gwrngs'
```

are described in vignettes.

The DataFrame function is imported from IRanges.

The SnpMatrix-class is used to represent data related to rule-based imputation, using the impute.snps function.

si.hs.38 is a Seqinfo-class instance for hg38.

nodeData (and nodes, ugraph, subGraph, adj) are exported for use in the vignettes.

Author(s)

VJ Carey <stvjc@channing.harvard.edu>

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References

```
http://www.genome.gov/gwastudies/.
```

Partial support from the Computational Biology Group at Genentech, Inc.

Examples

```
## Not run:
  data(ebicat38)
  ebicat38
## End(Not run)
```

bindcadd_snv

bind CADD scores of Kircher et al. 2014 to a GRanges instance

Description

bind CADD scores of Kircher et al. 2014 to a GRanges instance; by default will use HTTP access at UW

Usage

```
bindcadd_snv(gr, fn = "http://krishna.gs.washington.edu/download/CADD/v1.0/1000G.tsv.gz")
```

Arguments

gr query ranges to which CADD scores should be bound

fn path to Tabix-indexed bgzipped TSV of CADD as distributed at krishna.gs.washington.edu

on 1 April 2014

Details

joins CADD fields at addresses that match query; the CADD fields for query ranges that are not matched are set to NA

Value

GRanges instance with additional fields as obtained in the CADD resource

Note

This software developed in part with support from Genentech, Inc.

Author(s)

VJ Carey <stvjc@channing.harvard.edu>

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References

M Kircher, DM Witten, P Jain, BJ O'Roak, GM Cooper, J Shendure, A general framework for estimating the relative pathogenicity of human genetic variants, Nature Genetics Feb 2014, PMID 24487276

Examples

```
## Not run:
# requires internet access
data(ebicat37)
g2 = as(ebicat37, "GRanges")
bindcadd_snv( g2[which(seqnames(g2)=="chr2")][1:20] )
## End(Not run)
```

gwastagger

data on 1000 genomes SNPs that 'tag' GWAS catalog entries

Description

data on 1000 genomes SNPs that 'tag' GWAS catalog entries

Usage

```
data(gwastagger)
```

Format

```
The format is:
```

```
Formal class 'GRanges' [package "GenomicRanges"] with 6 slots
..@ segnames :Formal class 'Rle' [package "IRanges"] with 4 slots
.....@ values: Factor w/ 24 levels "chr1", "chr2", ..: 1 2 3 4 5 6 7 8 9 10 ...
.....@ lengths: int [1:22] 24042 23740 21522 14258 14972 34101 12330 11400 8680 15429 ...
.....@ elementMetadata: NULL
.. .. ..@ metadata : list()
..@ ranges :Formal class 'IRanges' [package "IRanges"] with 6 slots
.....@ start: int [1:297579] 986111 988364 992250 992402 995669 999686 1005579 1007450
1011209 1011446 ...
.....@ width: int [1:297579] 1 1 1 1 1 1 1 1 1 1 ...
.. .. ..@ 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 297579
.. .. ..@ elementMetadata: NULL
```

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```
.....@ metadata : list()
..@ elementMetadata:Formal class 'DataFrame' [package "IRanges"] with 6 slots
.....@ rownames: NULL
.....@ nrows: int 297579
.. .. .. @ listData :List of 3
..... $\tagid : \chr [1:297579] \"rs28479311" \"rs3813193" \"\chr1:992250" \"rs60442576" \...
..... $R2: num [1:297579] 0.938 0.994 0.969 1 1 ...
..... $\shatsa \text{baseid: chr [1:297579] "rs3934834" "rs3934834" "rs3934834" "rs3934834" ...
.....@ elementType : chr "ANY"
.....@ elementMetadata: NULL
.....@ metadata : list()
..@ seqinfo :Formal class 'Seqinfo' [package "GenomicRanges"] with 4 slots
.....@ seqnames : chr [1:24] "chr1" "chr2" "chr3" "chr4" ...
.....@ seqlengths: int [1:24] 249250621 243199373 198022430 191154276 180915260 171115067
159138663 146364022 141213431 135534747 ...
.....@ is_circular: logi [1:24] FALSE FALSE FALSE FALSE FALSE FALSE ...
.....@ genome: chr [1:24] "hg19" "hg19" "hg19" "hg19" ...
..@ metadata : list()
```

Details

This GRanges instance includes locations for 297000 1000 genomes SNP that have been identified as exhibiting LD with NHGRI GWAS SNP as of September 2013. The tagid field tells the name of the tagging SNP, the baseid field is the SNP identifier for the GWAS catalog entry, the R2 field tells the value of R-squared relating the distributions of the tagging SNP and the GWAS entry. Only tagging SNP with R-squared 0.8 or greater are included. A self-contained R-based procedure should emerge in 2014.

Source

NHGRI GWAS catalog; plink is used with the 1000 genomes VCF in a perl routine by Michael McGeachie, Harvard Medical School;

```
data(gwastagger)
gwastagger[1:5]
data(ebicat37)
mean(ebicat37$SNPS %in% gwastagger$baseid)
# ideally, all GWAS SNP would be in our tagging ranges as baseid
query <- setdiff(ebicat37$SNPS, gwastagger$baseid)
# relatively recent catalog additions
sort(table(ebicat37[which(ebicat37$SNPS %in% query)]$DATE.ADDED.TO.CATALOG), decreasing=TRUE)[1:10]</pre>
```

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gwaswloc-class

Class "gwaswloc"

Description

A container for GRanges instances representing information in the NHGRI GWAS catalog.

Objects from the Class

Objects can be created by calls of the form new("gwaswloc", ...). Any GRanges instance can be supplied.

Slots

```
extractDate: character set manually in .onAttach code to indicate date of retrieval of base table seqnames: Object of class "Rle" typically representing chromosome numbers of loci associated with specific traits

ranges: Object of class "IRanges" genomic coordinates for locus

strand: Object of class "Rle" identifier of chromosome strand

elementMetadata: Object of class "DataFrame" general DataFrame-class instance providing attributes for the locus-trait association

seqinfo: Object of class "Seqinfo"

metadata: Object of class "list"
```

Extends

```
Class "GRanges", directly. Class "GenomicRanges", by class "GRanges", distance 2. Class "Vector", by class "GRanges", distance 3. Class "GenomicRangesORmissing", by class "GRanges", distance 3. Class "GenomicRangesORGRangesList", by class "GRanges", distance 3. Class "Annotated", by class "GRanges", distance 4.
```

Methods

```
[ signature(x = "gwaswloc"): a character argument to the bracket will be assumed to be a db-
SNP identifier for a SNP locus, and records corresponding to this SNP are extracted; numeric
indexes are supported as for GRanges-class instances.
```

```
getRsids signature(x = "gwaswloc"): extract all dbSNP identifiers as a character vector
```

subsetByChromosome signature(x = "gwaswloc"): select records by chromosome, a vector
 of chromosomes may be supplied

subsetByTraits signature(x = "gwaswloc"): select all records corresponding to a given vector
 of traits

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Note

In gwascat package 1.9.6 and earlier, the globally accessible gwaswloc instance gwrngs was created upon attachment. This is no longer the case.

Author(s)

VJ Carey <stvjc@channing.harvard.edu>

References

```
http://www.genome.gov/gwastudies/
```

Examples

```
showClass("gwaswloc")
```

gwcex2gviz

Prepare salient components of GWAS catalog for rendering with Gviz

Description

Prepare salient components of GWAS catalog for rendering with Gviz

Usage

Arguments

basegr	gwaswloc instance containing information about GWAS in catalog	
contextGR	A GRanges instance delimiting the visualization in genomic coordinates	
txrefpk	a TxDb package, typically	
genesympk	string naming annotationDbi .db package	
genome	character tag like 'hg19'	
plot.it	logical, if FALSE, just return list	
maxmlp	maximum value of -10 log p $-$ winsorization of all larger values is performed, modifying the contents of Pvalue_mlogp in the elementMetadata for the call	

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Examples

```
args(gwcex2gviz)
#gwascat:::.onAttach("", "gwascat")
data(ebicat37)
seqlevelsStyle(ebicat37) = "UCSC"
gwcex2gviz(ebicat37)
```

gwdf_2012_02_02

internal data frame for NHGRI GWAS catalog

Description

convenience container for imported table from NHGRI GWAS catalog

Usage

```
data("gwdf_2014_09_08")
```

Format

A data frame with 17832 observations on the following 34 variables.

'Date Added to Catalog' a character vector

PUBMEDID a numeric vector

'First Author' a character vector

Date a character vector

Journal a character vector

Link a character vector

Study a character vector

'Disease/Trait' a character vector

'Initial Sample Size' a character vector

'Replication Sample Size' a character vector

Region a character vector

Chr_id a character vector

Chr_pos a character vector

'Reported Gene(s)' a character vector

Mapped_gene a character vector

Upstream_gene_id a character vector

Downstream_gene_id a character vector

Snp_gene_ids a character vector

Upstream_gene_distance a character vector

Downstream_gene_distance a character vector

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```
'Strongest SNP-Risk Allele' a character vector
SNPs a character vector

Merged a character vector
Snp_id_current a character vector
Context a character vector
Intergenic a character vector
'Risk Allele Frequency' a character vector
'p-Value' a character vector
Pvalue_mlog a character vector
'p-Value (text)' a character vector
'OR or beta' a character vector
'95% CI (text)' a character vector
'Platform [SNPs passing QC]' a character vector
CNV a character vector
```

Note

In versions prior to 1.9.6, The .onAttach function specifies which data frame is transformed to GRanges. This is now managed manually.

Source

```
http://www.genome.gov/gwastudies
```

Examples

```
## Not run:
data(gwdf_2014_09_08)
# try gwascat:::gwdf2GRanges on this data.frame
## End(Not run)
```

ldtagr

expand a list of variants by including those in a VCF with LD exceeding some threshold

Description

expand a list of variants by including those in a VCF with LD exceeding some threshold

Usage

```
ldtagr(snprng, tf, samples, genome = "hg19", lbmaf = 0.05, lbR2 = 0.8, radius = 1e+05)
```

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Arguments

snprng	a named GRanges for a single SNP. The name must correspond to the name that will be assigned by <code>genotypeToSnpMatrix</code> to the corresponding column of a SnpMatrix.
tf	TabixFile instance pointing to a bgzipped tabix-indexed VCF file
samples	a vector of sample identifiers, if excluded, all samples used
genome	tag like 'hg19'
lbmaf	lower bound on variant MAF to allow consideration
1bR2	lower bound on R squared for regarding SNP to be incorporated

radius of search in bp around the input range

Details

radius

```
uses snpStats ld()
```

Value

a GRanges with names corresponding to 'new' variants and mcols fields 'paramRangeID' (base variant input) and 'R2'

Note

slow but safe approach. probably a matrix method could be substituted using the nice sparse approach already in snpStats

Author(s)

VJ Carey

```
require(GenomicRanges)
cand = GRanges("1", IRanges(113038694, width=1))
names(cand) = "rs883593"
require(VariantAnnotation)
expath = dir(system.file("vcf", package="GGtools"), patt=".*exon.*gz$", full=TRUE)
tf = TabixFile(expath)
ldtagr( cand, tf, lbR2 = .8)
# should do with 1000 genomes in S3 bucket and gwascat
```

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locon6

location information for 10000 SNPs probed on Affy GW 6.0

Description

location information for 10000 SNPs probed on Affy GW 6.0

Usage

```
data(locon6)
```

Format

A data frame with 10000 observations on the following 3 variables.

```
dbsnp_rs_id a character vector
chrom a character vector
physical_pos a numeric vector
```

Details

extracted from pd.genomewidesnp.6 v 1.4.0; for demonstration purposes

Examples

```
data(locon6)
str(locon6)
```

makeCurrentGwascat

read NHGRI GWAS catalog table and construct associated GRanges instance

Description

read NHGRI table and construct associated GRanges instance

Usage

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Arguments

table.url string identifying the .txt file curated at EBI/EMBL

fixNonASCII logical, if TRUE, non-ASCII characters as identified by iconv will be replaced

by asterisk

genome character string: 'GRCh38' is default and yields current image as provided by

EMBL/EBI; 'GRCh37' yields a realtime liftOver to hg19 coordinates, via An-

notationHub storage of the chain files. Any other value yields an error.

withOnt logical indicating whether 'alternative' (ontology-present, includes repetition of

loci with one:many ontological mapping) or 'full' (ontology-absent, one record

per locus report) version of distributed table

Details

records for which clear genomic position cannot be determined are dropped from the ranges instance an effort is made to use reasonable data types for GRanges metadata, so some qualifying characters such as (EA) in Risk allele frequency field will simply be omitted during coercion of contents of that field to numeric.

Value

a GRanges instance

Author(s)

VJ Carey

Examples

```
## Not run:
# if you have good internet access
newcatr = makeCurrentGwascat()
## End(Not run)
```

obo2graphNEL

convert a typical OBO text file to a graphNEL instance (using Term elements)

Description

convert a typical OBO text file to a graphNEL instance (using Term elements)

Usage

```
obo2graphNEL(obo, kill = "\\[Typedef\\]", killTrailSp=TRUE)
node2uri(nn)
uri2node(us)
```

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Arguments

obo	string naming a file in OBO format
nn	node name for converted graph, generally of form EFO:nnnnnnn
us	URI string from GWAS catalog annotation MAPPED_TRAIT_URI
kill	entity types to be excluded from processing – probably this should be in a 'keep' form, but for now this works.
killTrailSp	In the textual version of EFO ca. Aug 2015, there is a trailing blank in the tag field defining EFO:0000001, which is not present in references to this term. Set this to TRUE to eliminate this, or graphNEL construction will fail to validate.

Details

Very rudimentary list and grep operations are used to retain sufficient information to map the DAG to a graphNEL, using formal term identifiers as node names and 'is-a' relationships as edges, and term names and other metadata are assigned to nodeData components.

Value

```
a graphNEL instance
```

Note

The OBO for Human Disease ontology is serialized as text with this package.

Author(s)

VJ Carey <stvjc@channing.harvard.edu>

References

For use with human disease ontology, http://www.obofoundry.org/cgi-bin/detail.cgi?id=disease_ontology

```
data(efo.obo.g)
hn = nodes(efo.obo.g)[1:5]
hn
nodeData(efo.obo.g, hn[5])
```

14 riskyAlleleCount

ricky	ما ۱ ۵۰	leCount

given a matrix of subjects x SNP calls, count number of risky alleles

Description

given a matrix of subjects x SNP calls, count number of risky alleles for various conditions, relative to NHGRI GWAS catalog

Usage

```
riskyAlleleCount(callmat, matIsAB = TRUE, chr,
  gwwl , snpap = "SNPlocs.Hsapiens.dbSNP.20120608",
  gencode = c("A/A", "A/B", "B/B"))
```

Arguments

callmat	matrix with subjects as rows, SNPs as columns; entries can be generic A/A, A/B, B/B, or specific nucleotide calls
matIsAB	logical, FALSE if nucleotide codes are present, TRUE if generic call codes are present; in the latter case, gwascat:::ABmat2nuc will be run
chr	code for chromosome, should work with the SNP annotation getSNPlocs function, so likely " $ch[nn]$ "
gwwl	an instance of gwaswloc
snpap	name of a Bioconductor SNPlocs. Hsapiens.dbSNP.* package
gencode	codes used for generic SNP call

Value

matrix with rows corresponding to subjects, columns corresponding to SNP

```
data(gg17N) # translated from GGdata chr 17 calls using ABmat2nuc
data(ebicat37)
library(GenomeInfoDb)
seqlevelsStyle(ebicat37) = "UCSC"
h17 = riskyAlleleCount(gg17N, matIsAB=FALSE, chr="ch17", gwwl=ebicat37)
h17[1:5,1:5]
table(as.numeric(h17))
```

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GWAS catalog

Description

```
operations on GWAS catalog
```

Usage

```
topTraits (gwwl, n=10, tag="DISEASE.TRAIT")
locs4trait(gwwl, trait, tag="DISEASE.TRAIT")
chklocs(chrtag="20", gwwl)
```

Arguments

gwwl	instance of gwaswloc
n	numeric, number of traits to report
tag	character, name of field to be used for trait enumeration
trait	character, trait to use for filtering
chrtag	character, chromosome identifier

Value

topTraits returns a character vector of most frequently occurring traits in the database locs4trait returns a gwaswloc object with records defining associations to the specified trait chklocs returns a logical that is TRUE when the asserted locations of SNP in the GWAS catalog agree with the locations given in the dbSNP package SNPlocs.Hsapiens.dbSNP.20110815

Author(s)

VJ Carey <stvjc@channing.harvard.edu>

```
data(ebicat38)
topTraits(ebicat38)
```

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	traitsManh	use ggbio facilities to display GWAS results for selected traits in genomic coordinates
--	------------	---

Description

use ggbio facilities to display GWAS results for selected traits in genomic coordinates

Usage

```
traits Manh (gwr, selr = GRanges (seqnames = "chr17", IRanges (3e+07, 5e+07)), traits = c("Asthma", "Park (seqnames = "chr17", IRanges (seqnames = "chr17")), traits = c("Asthma", "Park")
```

Arguments

gwr	GRanges instance as managed by the gwaswloc container design, with Disease.Trait and Pvalue_mlog among elementMetadata columns
selr	A GRanges instance to restrict the gwr for visualization. Not tested for noncontiguous regions.
traits	Character vector of traits to be exhibited; GWAS results with traits not among these will be labeled "other".
truncmlp	Maximum value of $-\log 10$ p to be displayed; in the raw data this ranges to the hundreds and can cause bad compression.
	not currently used

Details

uses a ggbio autoplot

Value

autoplot value

Author(s)

VJ Carey <stvjc@channing.harvard.edu>

```
# do a p-value truncation if you want to reduce compression
data(ebicat38)
library(GenomeInfoDb)
seqlevelsStyle(ebicat38) = "UCSC"
traitsManh(ebicat38)
```

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