# GGBase

# October 5, 2010

GGbase-package GGbase Package Overview

## Description

GGbase Package Overview

# Details

This package provides infrastructure for programming related to the genetics of gene expression. The GGtools package makes use of classes and methods defined in this package. GGdata and hmyriB36 packages use the class structures defined in this package for serialized data.

Introductory information is available from vignettes, type openVignette().

Full listing of documented man pages is available in HTML view by typing help.start() and selecting GGbase package from the Packages menu or via library (help="GGbase").

# Author(s)

V. Carey

MAFfilter

restrict SNP in an smlSet to range of minor allele frequencies (MAF)

## Description

restrict SNP in an smlSet to range of minor allele frequencies (MAF)

## Usage

MAFfilter(x, lower = 0, upper = 1)

## Arguments

Х	smlSet instance
lower	numeric lower bound on minor allele frequency for keeping a SNP
upper	numeric upper bound on minor allele frequency for keeping a SNP

## Details

uses snp.matrix-class summary method from snpMatrix

# Value

revised instance of smlSet-class

## Author(s)

VJ Carey <stvjc@channing.harvard.edu>

## Examples

```
data(smlSet.example)
sapply(smList(MAFfilter(smlSet.example, lower=.1)), dim)
```

SessionInfo-class Class "SessionInfo" – objects to help stamp an output with information on session state

## Description

Class "SessionInfo" - objects to help stamp an output with information on session state

## **Objects from the Class**

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

# Slots

.S3Class: Object of class "character" simple cast to allow checking

## Extends

Class "oldClass", by class "sessionInfo", distance 2.

#### Methods

No methods defined with class "SessionInfo" in the signature.

## Examples

showClass("SessionInfo")

genesym-class Class "genesym" and other casting classes

#### Description

classes that help establish symbol semantics for dispatching

#### **Objects from the Class**

Objects can be created by calls of the form new ("genesym", ...), or by special constructor functions. As of GGBase version 3.7.1, you can use genesym(...), chrnum(...), probeld(...), rsid(...). These generally just extend character or numeric so that vector operations are straightforward, but attach type information so that methods such as [ 'know' what they are getting.

Currently, genesym is used to allow HUGO symbols to be passed to [; chrnum identifies numerals or numeric constants as indices into the set of chromosomes (no chr prefix is allowed); rsid identifies dbSNP identifiers; probeId identifies a string as a microarray probe identifier.

snpdepth identifies a number that will be used as the number of chromosome-specific test results to be retained in any genome-wide screen

# Slots

.Data: Object of class "character" ~~

## Extends

Class "character", from data part. Class "vector", by class "character", distance 2. Class characterORMIAME, by class "character", distance 2.

## Author(s)

VJ Carey <stvjc@channing.harvard.edu>

## Examples

```
showClass("genesym")
genesym("CPNE1")
```

featureFilter remove unannotated or undesired features from an smlSet instance

## Description

remove unannotated or undesired features from an smlSet instance

# Usage

```
featureFilter(x, requires = c("loc", "autosomal"))
```

#### Arguments

Х	instance of smlSet class
requires	character vector – if "loc" is present, require that a non-NA value is present in CHRLOC for each feature; if "autosomal" is present, require that CHR value is in 1:22 (presently assumes human genome)

## Value

revised smlSet instance excluding features no

## Author(s)

VJ Carey

#### Examples

```
data(smlSet.example)
dim(exprs(smlSet.example))
fff = featureFilter(smlSet.example)
dim(exprs(fff))
```

```
gwSnpScreenResult-class
```

Class "gwSnpScreenResult" – containers for GGtools gwSnpScreen method outputs and allied objects

#### Description

Class "gwSnpScreenResult" - container for GGtools gwSnpScreen method outputs and allied objects

#### **Objects from the Class**

Objects can be created by calls of the form new ("gwSnpScreenResult", ...). These will be primarily lists of inference tables (snps are rows, columns are statistics and p-values). Additional slots manage analysis metadata.

gwSnpScreenResult is intended for genome-wide analysis of expression for a single gene.

cwSnpScreenResult is intended for the restriction to a single chromosome.

multiGwSnpScreenResult is intended for analyses with multiple genes.

Because the vast majority of tests are uninformative, early filtering is important for managing object sizes. Instances of filteredGwSnpScreenResult and filteredMultiGwSnpScreenResult are created when a snpdepth parameter is used with gwSnpTests.

#### make\_smlSet

## Slots

- .Data: Object of class "list" containing inference tables (snps are rows, columns are statistics and p-values)
- gene: Object of class "character" typically the HUGO symbol of the gene analyzed
- psid: Object of class "character" the feature identifier of the associated microarray
- annotation: Object of class "character" vector of relevant annotation package identifier names
- formula: Object of class "formula" the formula used to fit the model relating expression to genotype

#### Extends

Class "list", from data part. Class "vector", by class "list", distance 2. Class AssayData, by class "list", distance 2.

# Methods

plot and show

# Author(s)

VJ Carey <stvjc@channing.harvard.edu>

## Examples

```
showClass("gwSnpScreenResult")
showClass("cwSnpScreenResult")
```

make\_smlSet create an smlSet instance from components

## Description

create an smlSet instance from components

## Usage

```
make_smlSet(es, sml, organism = "Homo sapiens")
```

#### Arguments

es	ExpressionSet instance
sml	list of snp.matrix instances
organism	string naming organism

## Details

combines snp.matrix instances with expression data

## Value

instance of smlSet class

# Author(s)

VJ Carey <stvjc@channing.harvard.edu>

#### Examples

```
data(smlSet.example) # here we just show the mechanics from a working smlSet
es = as(smlSet.example, "ExpressionSet")
sl = smList(smlSet.example)
mm = make_smlSet(es, sl)
validObject(mm)
mm
```

# Description

object to contain results of restricted gene-centric searches for eQTL

#### **Objects from the Class**

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

## Slots

```
.Data: Object of class "list" - list of results of snp.rhs.tests
conditions: Object of class "list" - list of runtime conditions encountered
call: Object of class "call" - for auditing, the call used is saved
```

### Extends

```
Class "list", from data part. Class "vector", by class "list", distance 2. Class "AssayData", by class "list", distance 2.
```

#### Methods

```
show signature(object = "multiCisTestResult"):...
```

#### Author(s)

VJ Carey <stvjc@channing.harvard.edu>

## Examples

showClass("multiCisTestResult")

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plot\_EvG-methods formal method for visualizing expression distributions vs genotype

## Description

boxplot expression vs genotype

#### Methods

```
gsym = "genesym", rsid = "rsid", sms = "smlSet" generates an annotated boxplot
multisnp methods plot\_EvG2 allows specification of a second SNP rsid and shows boxplots over
```

the cross-tabulation of the allele combinations

## Examples

```
data(smlSet.example)
plot_EvG(genesym("WBP5"), rsid("rs10483083"), smlSet.example)
```

smlSet-class	Documentation on S4 class "smlSet" an eSet-derived container for
	snpMatrix lists, allowing efficient combination of SNP chip genotyping
	with microarray expression data, and allied classes

#### Description

Documentation on S4 class "smlSet" an eSet-derived container for snpMatrix lists, allowing efficient combination of SNP chip genotyping with microarray expression data, and allied classes

## **Objects from the Class**

Objects can be created by calls of the form new("smlSet", assayData, phenoData, featureData, experimentData, annotation, ...). These objects respond to interrogation on samples, expression values, SNP values, and other metadata.

## Slots

- organism: Object of class "character" informal, "Hs" recommended for human
- assayData: Object of class "AssayData" intended to hold expression data coordinated with the smlEnv data
- phenoData: Object of class "AnnotatedDataFrame" standard sample-level data container from eSet design
- featureData: Object of class "AnnotatedDataFrame" standard feature-level metadata container, implied usage is for documenting the expression data elements
- annotation: Object of class "character" vector giving the Bioconductor annotation package (.db type) for decoding expression feature identifiers.
- .\_\_\_classVersion\_\_: Object of class "Versions" class version tracking metadata

#### Extends

Class eSet, directly. Class VersionedBiobase, by class "eSet", distance 2. Class Versioned, by class "eSet", distance 3.

# Methods

smList signature(x = "smlSet"): retrieves the actual list of snp.matrix entities
smlEnv signature(x = "smlSet"): retrieves the environment holding snp.matrix entities

- **exprs** signature (x = "smlSet"): retrieves the matrix of expression values
- snps signature(x = "smlSet", chr = "chrnum"): retrieves the raw matrix of genotype values (snp.matrix instance from snpMatrix package

combine: concatenates expression data and forms intersection of SNP sets

```
getAlleles (smlSet, rsid): returns A/B notations for SNP determined by rsid
```

coerce: extracts exprs, phenoData and annotation and constructs ExpressionSet

[ signature(x = "smlSet", i = "ANY", j = "ANY", drop = "ANY"): Quick methods for subsetting elements of smlSets have been provided. If X is an smlSet instance and G is a vector of class probeId-class, then X[G,] will reduce the expression data to the probes specified in G. If X is an smlSet instance and G is a vector of class chrnum-class, then X[G,] will reduce the SNP genotype data to the SNPs resident on chromosomes enumerated in G.

If X is an smlSet instance and G is a vector of class rsid-class, then X[G,] will reduce the SNP genotype data to the SNPs enumerated in the dbSNP id in G.

#### Note

We have included a [method for snp.matrix instances that accepts an rsid instance as a column selector.

#### Author(s)

VJ Carey <stvjc@channing.harvard.edu>

## See Also

GGtools package makes extensive use of these classes and methods.

```
showClass("smlSet")
data(smlSet.example)
smlSet.example
validObject(smlSet.example)
# workout on expression components
dim(exprs(smlSet.example))
fn = featureNames(smlSet.example)[1:10]
fn
ss2 = smlSet.example[ probeId(fn), ] # restrict exprs to set of probes
dim(exprs(ss2))
# workout on SNP components
smList(smlSet.example)
dim(smList(ss2)[[1]])
ss2[ chrnum(21), ] # trivial restriction of SNP to a chromosome
```

#### smlSummary

```
sn = colnames(smList(ss2)[[1]])[1:20] # get some dbSNP ids
ss3 = ss2[ rsid(sn), ] # subset the snps
dim(smList(ss3)[[1]])
dim(smList(ss3)[["21"]]) # check names
ss3
as(snps(ss3, chrnum(21)), "character")[1:5,1:5] # generic codes
as(snps(ss3, chrnum(21)), "numeric")[1:5,1:5] # number copies of B
as(snps(ss3, chrnum(21)), "matrix")[1:5,1:5] # raw
```

smlSummary

class and function to summarize frequency information on genotypes in an smlSet

## Description

generates information on sample size, minor allele frequency, specific call frequencies, and HWE test results on all SNP in an smlSet

## Usage

smlSummary(x)

## Arguments

x instance of smlSet-class

## Details

to control volume of printout a simple list extending class is defined for show method

# Value

Instance of smlSummary class, which simply extends list. Each list element is a matrix of results provided by summary, snp.matrix-method.

#### Author(s)

VJCarey <stvjc@channing.harvard.edu>

```
data(smlSet.example)
smlSummary(smlSet.example)
```

snpLocs.Hs

#### Description

SNP location accessor

## Usage

snpLocs.Hs(cnum, rsid)

#### Arguments

cnum	chrnum instance indicating chromosome
rsid	rsid instance giving dbSNP ids for snps of interest

#### Details

The SNPlocs.Hsapiens.dbSNP.\* package is curated by Bioconductor and maintains location and allele information on SNP.

The hsSnpLocs environment is available through data(hsSnpLocs), and contains a unified representation of the information in the SNPlocs package (which is only available through various chromosome-specific calls).

## Value

two-row matrix – top row is numeric suffix of dbSNP ids, bottom row is location, genome-wide if chrnum is missing

## Author(s)

Vince Carey <stvjc@channing.harvard.edu>

```
data(smlSet.example)
nn21 = colnames(smList(smlSet.example)[["21"]])
length(nn21)
ss = snpLocs.Hs(chrnum(21), rsid(nn21))
dim(ss)
# notes that the locations available are not completely
# congruent with those asserted in HapMap data
```

snpsNear

## Description

obtain list of rs numbers for snps near a gene

## Usage

```
snpsNear(sym, radius=1e+05, chrnum, ...)
```

#### Arguments

sym	instance of genesym class [e.g., use genesym(string) for gene 'string'], or of
	rsid class, or of numeric class. An instance of GeneSet-class can also be
	supplied if it has geneIdType AnnotationIdentifier.
radius	number of base-pairs in each direction to look
chrnum	chrnum instance optional
	options not now in use

## Details

simple arithmetic based on output of snpLocs.Hs

#### Value

character vector of rsxxxxx, dbSNP id, according to locations from SNPlocs.Hsapiens.dbSNP.20071016 package, as transferred to snpLocs.Hs resource in GGBase

note that an attribute 'target' is returned, a named vector with components chr and loc describing chromosome and location of the target for which nearby SNPs are sought

## Note

first invocation can take longer than subsequent, if snpLocs.Hs has not been invoked previously

## Author(s)

Vince Carey <stvjc@channing.harvard.edu>

```
nearc = snpsNear(genesym("BACH1"), 10000, chrnum(21))
data(smlSet.example)
ss = smList(smlSet.example)[[1]]
# following calculation requires new "[" for j an instance of rsid
clo = ss[, rsid(snpsNear(rsid("rs6060535"), rad=1500, chrnum(20))) ]
clo
# try a gene set
library(GSEABase)
s1 = GeneSet(c("BACH1", "ATP50"), geneIdType=SymbolIdentifier())
s2 = s1
geneIdType(s2) = AnnotationIdentifier("illuminaHumanv1.db")
```

snpsNear

```
s2
sapply(snpsNear(s2), length)
```

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