# Package 'GGBase'

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Author VJ Carey <stvjc@channing.harvard.edu></stvjc@channing.harvard.edu>	
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Maintainer VJ Carey <stvjc@channing.harvard.edu></stvjc@channing.harvard.edu>	
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transformations of expression data in smlSet instances

# **Description**

transformations of expression data in smlSet instances or assay data in SummarizedExperiment

# Usage

```
clipPCs(x, inds2drop, center = TRUE)
regressOut(sms, rhs, ...)
```

# **Arguments**

x instance of smlSet or RangedSummarizedExperiment sms instance of smlSet or RangedSummarizedExperiment

inds2drop 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 inds2drop, the smlSet is returned unchanged, with a message.

center logical, passed to prcomp

rhs formula fragment (no dependent variable) used to form residuals in a reexpres-

sion of the expression matrix; variable bindings found in pData of an Expres-

sionSet or colData of a SummarizedExperiment

... arguments passed to lmFit

#### **Details**

clipPCs is an operation on the n x 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

```
an instance of smlSet
```

#### 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. The PCA-based method seems to have reasonable effectiveness in examples worked with GGdata.

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#### **Examples**

externalize

create a package with a decomposed smlSet instance from a unified smlSet instance, to reduce memory footprints

#### **Description**

create a package with a decomposed smlSet instance from a unified smlSet instance, to reduce memory footprints

# Usage

```
externalize(smlSet, packname,
   author = "Replace Me <auth@a.b.com>",
   maintainer = "Replace Me <repl@a.b.com>")
allsnps(packname)
```

## **Arguments**

smlSet an smlSet instance to be regarded as a source or template for a package that can

be used with greater efficiency of access to genotype contents

packname name of the package to be generated

author string to be used in DESCRIPTION file of generated package maintainer string to be used in DESCRIPTION file of generated package

# **Details**

The genotype content of the input smlSet is separated into separate RDA files in the inst/parts folder of the package to be generated. The ExpressionSet element of the input smlSet is stored as object ex in file eset.rda in the data folder of the package to be generated.

# Value

creates folder structure and metadata for an installable R package

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#### Note

allsnps() gives the list of vectors of snpnames in use in the package

#### Author(s)

VJ Carey <stvjc@channing.harvard.edu>

#### **Examples**

```
## Not run:
setwd(tempdir())
sms = getSS("GGtools", c("20", "21"))
externalize(sms, "demopack")
dir()
dir("demopack")
## End(Not run)
```

MAFfilter

Filter genotype contents of an smlSet according to certain SNP allele frequency features.

# Description

Filter genotype contents of an smlSet according to certain SNP allele frequency features.

# Usage

```
MAFfilter(x, lower = 0, upper = 1)
GTFfilter(x, lower = 0)
dropMonomorphies(sms)
```

# **Arguments**

```
x smlSet-class instance
sms smlSet-class instance
```

lower lower bound on MAF or GTF to allow retention of associated locus upper upper bound on MAF or GTF to allow retention of associated locus

# **Details**

uses col. summary to compute MAF or GTF. dropMonomorphies also uses col.summary.

#### Value

```
smlSet-class instance
```

#### Author(s)

VJ Carey <stvjc@channing.harvard.edu>

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#### **Examples**

```
if (file.exists(system.file("parts/20.rda", package="GGtools"))) {
c20 = getSS("GGtools", "20")
c20f = MAFfilter(c20, lower=.05)
c20f
```

 ${\tt make\_smlSet}$ 

construct an smlSet instance from existing resources

# **Description**

construct an smlSet instance from existing resources, either using ExpressionSet and SnpMatrix instances, or a suitably structured package

# Usage

```
make_smlSet(es, sml, organism = "Homo sapiens", harmonizeSamples = FALSE)
getSS(packname, chrs, renameChrs, probesToKeep=NULL, exFilter=function(x)x,
   wrapperEndo=NULL, checkValid=TRUE)
```

#### Arguments

instance of ExpressionSet-class es sml named list of SnpMatrix-class instances organism conventional token for species harmonizeSamples

logical indicating whether steps should be taken to be sure that the components represent identical sets of samples. can be time consuming so defaults to FALSE, in which case the user must be sure that the genotype and expression

components are compatible in terms of sample content

packname string naming the installed package from which expression and genotype data

will be acquired

chrs character vector naming the prefixes of genotype files to be used in the resulting

smlSet instance; see notes below

character vector of same length as chrs specifying one-to-one renaming operarenameChrs

tion for genotype components in smlSet

probesToKeep character vector identifying probes to be retained in the constructed smlSet – to

be deprecated in favor of exFilter

exFilter function that should accept and return ExpressionSet, will be executed just

after probesToKeep filter if present

wrapperEndo function that accepts and returns an smlSet instance, allowing any sort of trans-

formation of contents acquired with getSS, executed after all assembly and fil-

tering completed.

checkValid logical, if TRUE, function will fail if created smlSet instance does not pass vali-

dObject()

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#### **Details**

Packages that work with getSS can be created out of existing smlSet instances using externalize.

#### Value

Instance of smlSet-class.

# Author(s)

VJ Carey <stvjc@channing.harvard.edu>

# **Examples**

```
if ("GGtools" %in% installed.packages()[,1]) {
   s20 = getSS("GGtools", "20", renameChrs="chr20")
   s20
   make_smlSet( as(s20, "ExpressionSet"), smList(s20) )
}
```

plot\_EvG

display the association between expression values and genotypes in an smlSet instance

# **Description**

display the association between expression values and genotypes in an smlSet instance

# Usage

```
plot_EvG(gsym, rsid, sms, ...)
```

#### **Arguments**

gsym	instance of class genesym or probeId, casting a string that names a gene (which will be looked up using the annotation slot of sms) or a probe which must be resident on the array underlying the expression content of sms
rsid	instance of class rsid naming a SNP with genotype values given among the columns of the smList components of sms
sms	an instance of smlSet
	additional parameter to plot

# **Details**

When the genotype is categorical, will use boxplots; when genotype has been imputed and includes expected allele counts, will use a scatterplot.

# Value

a plot is rendered on the current display

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#### Author(s)

VJ Carey <stvjc@channing.harvard.edu>

# **Examples**

```
if ("GGtools" %in% installed.packages()[,1]) {
  s20 = getSS("GGtools", "20")
  plot_EvG(genesym("CPNE1"), rsid("rs6060535"), s20)
}
```

rsid-class

Class "rsid"

# **Description**

rsid(), probeId(), and genesym() are basic casting methods that assign a type to a token.

# **Objects from the Class**

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

#### **Slots**

.Data: Object of class "character" that holds the content to which a type is associated by the method.

# Extends

Class "character", from data part. Class "vector", by class "character", distance 2. Class "data.frameRowLabels", by class "character", distance 2. Class "SuperClassMethod", by class "character", distance 2. Class "character", distance 2. Class "character", distance 2. Class "character", distance 2. Class "atomicVector", by class "character", distance 2. Class "index", by class "character", distance 2.

# Methods

No methods defined with class "probeId" or "genesym" in the signature. However [ with a smlSet-class argument will dispatch differently if instances of these classes are supplied. Various GGtools reporting functions will use "rsid" for dispatch.

# Author(s)

VJ Carey <stvjc@channing.harvard.edu>

#### **Examples**

```
showClass("rsid")
```

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smlSet-class

Class "smlSet"

# **Description**

Integrative container for expression plus genotype data. Genotypes are stored in an efficient format defined in the snpStats package.

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

Objects can be created by calls of the form new("smlSet", assayData, phenoData, featureData, experimentData, an The make\_smlSet function can also be used to build smlSet instances.

#### **Slots**

```
smlEnv: Object of class "environment" that has a key smList element, to which a list of SnpMatrix-class instances is bound.
```

annotation: Object of class "character", describes featureNames component in terms of the name of the annotation package that can be used to decode expression probe names.

organism: Object of class "character", a conventional string.

assayData: Object of class "AssayData", manages the expression data. See AssayData-class.

phenoData: Object of class "AnnotatedDataFrame", manages sample level data. See AnnotatedDataFrame-class.

featureData: Object of class "AnnotatedDataFrame", manages metadata on expression probes.

experimentData: Object of class "MIAxE", manages metadata on experiment as a whole. See MIAxE-class.

protocolData: Object of class "AnnotatedDataFrame", additional storage for experimental protocol description. See eSet-class.

.\_\_classVersion\_: Object of class "Versions"; internal management of class version.

#### **Extends**

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

#### Methods

```
[ signature(x = "smlSet", i = "ANY", j = "ANY", drop = "ANY"): will restrict the content of the smlSet instance according to features of the arguments supplied. If x is numeric or a probeId-class, the expression content will be restricted. If y is numeric or character, samples will be restricted accordingly.
```

**combine** signature(x = "smlSet", y = "smlSet"): This method attempts to amalgamate two smlSet instances in the appropriate way – assuming that samples are disjoint.

nsFilter signature(eset = "smlSet"): executes genefilter's method (see nsFilter, ExpressionSet-method)
 on the expression content, and then propagates the additional genotype and sample level content unchanged.

smList signature(x = "smlSet"): retrieves the list of SnpMatrix instances defining the genotype
content.

exprs signature(x = "smlSet"): retrieves the matrix of expression values

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permEx signature(sms = "smlSet"): uses sample() to generate a permutation of sample indices so that expression data are permuted against genotype data, but original unpermuted
sample identifiers are preserved; this is necessitated by the rematching behavior of snp.rhs.tests.

# Author(s)

VJ Carey <stvjc@channing.harvard.edu>

# **Examples**

showClass("smlSet")

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