

# Package ‘ggtut’

March 25, 2013

**Title** support for tutorial on genetics of gene expression ISMB 2011

**Version** 0.0.43

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

**Description** various resources for genetics of expression with R/bioc

**Suggests** rtracklayer, GenomicFeatures, ChIPpeakAnno, TxDb.Hsapiens.UCSC.hg18.knownGene

**Depends** R (>= 2.14.0), GGtools (>= 3.11.32), ff, GenomicRanges, snpStats, GGdata, GenomicFeatures, ChIPpeakAnno, Rsamtools(>= 1.5.35), cheung2010, SNPLocs.Hsapiens.dbSNP.20120608, hmyriB36

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

**License** Artistic-2.0

**LazyLoad** yes

## R topics documented:

|                         |   |
|-------------------------|---|
| ggtut-package . . . . . | 1 |
| c17imp . . . . .        | 2 |
| g17rngsnr . . . . .     | 3 |
| observed17ceu . . . . . | 4 |
| rules.n43 . . . . .     | 5 |
| snpgr17 . . . . .       | 6 |

**Index**

7

---

ggtut-package

*support for tutorial on genetics of gene expression ISMB 2011*

---

## Description

various resources for genetics of expression with R/bioc

## Details

## Package:

## Version:

Suggests:

Depends:

**License:**

LazyLoa

Built:

CPS17 is a

`CS17` is a `CST17` scores instance, as is `PERCENT17`, serialized to reduce vignette computation time. `SVA11` is an `sva()` object.

## Index:

obtained from the International Haplotype Map Project (HapMap) [11] and the Human Genome Project (HGDP) [12].

hg18tx a function that returns a reference to a transcriptDb instance for hg18

getFixedBFL retrieve a BamFileList of 41 small RNA seq extracts with nicely formatted HapMap names

Further information is available in the following vignettes:

tut11 (source, pdf)

## Author(s)

VJ Carey <stvjc@channing.harvard.edu>

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

## Examples

```
library(ggtut)
o17 = observed17ceu()
o17
ch41 = getFixedBFL()
ch41
```

c17imp

*SnpMatrix instance with imputations from 1KG to CEU chr 17*

## Description

SnpMatrix instance with imputations from 1KG to CEU chr 17

## Usage

```
data(c17imp)
```

## Format

The format is:

```
Formal class 'SnpMatrix' [package "snpStats"] with 1 slots
..@ .Data: raw [1:90, 1:175170] 77 d6 77 2d ...
.. ..- attr(*, "dimnames")=List of 2
.. .. ..$ : chr [1:90] "NA06985" "NA06991" "NA06993" "NA06994" ...
.. .. ..$ : chr [1:175170] "chr17:1869" "rs17055023" "rs6565733" "rs34663111" ...
```

## Source

uses rules.n43

## Examples

```
library(snpStats)
data(c17imp)
c17imp
```

g17rngsnr

*ranges of genes on chrom 17 (uses hg18)*

## Description

ranges of genes on chrom 17 (uses hg18)

## Usage

```
data(g17rngsnr)
```

## Format

The format is:

```
Formal class 'GRanges' [package "GenomicRanges"] with 6 slots
..@ seqnames :Formal class 'Rle' [package "IRanges"] with 4 slots
... ..@ values : Factor w/ 1 level "chr17": 1
... ..@ lengths : int 475
... ..@ elementMetadata: NULL
... ..@ metadata : list()
..@ ranges :Formal class 'IRanges' [package "IRanges"] with 6 slots
... ..@ start : int [1:475] 39509647 50333051 46294586 77439016 38229969 37098653 45133689
58981554 44263371 17349602 ...
... ..@ width : int [1:475] 46894 61277 5753 3743 19335 2772 6839 43820 33858 86118 ...
... ..@ NAMES : chr [1:475] "GI_21237796-A" "GI_4885638-S" "GI_22035666-S" "GI_17572809-
S" ...
... ..@ elementType : chr "integer"
... ..@ elementMetadata: NULL
... ..@ metadata : list()
..@ strand :Formal class 'Rle' [package "IRanges"] with 4 slots
... ..@ values : Factor w/ 3 levels "+", "-", "*": 3
... ..@ lengths : int 475
```

```

... .. ..@ elementMetadata: NULL
... .. ..@ metadata : list()
..@ elementMetadata:Formal class 'DataFrame' [package "IRanges"] with 6 slots
... .. ..@ rownames : NULL
... .. ..@ nrows : int 475
... .. ..@ listData :List of 1
... .. ..$ probeid: chr [1:475] "GI_21237796-A" "GI_4885638-S" "GI_22035666-S" "GI_17572809-
S" ...
... .. ..@ elementType : chr "ANY"
... .. ..@ elementMetadata: NULL
... .. ..@ metadata : list()
..@ seqinfo :Formal class 'Seqinfo' [package "GenomicRanges"] with 3 slots
... .. ..@ seqnames : chr "chr17"
... .. ..@ seqlengths : int NA
... .. ..@ is_circular: logi NA
..@ metadata : list()

```

## Examples

```

data(g17rngsnr)
g17rngsnr
## maybe str(g17rngsnr) ; plot(g17rngsnr) ...

```

observed17ceu

*obtain access to ff-based archives of eQTL test results*

## Description

obtain access to ff-based archives of eQTL test results

## Usage

```

observed17ceu()
onePerm17ceu()

```

## Details

The underlying ff data were obtained as follows

```

dropMonomorphies = function(sms) { sl = smList(sms) summs = lapply(sl, col.summary)
todrop = lapply(summs, function(x) which(x[, "RAF"]==1 | x[, "RAF"]==0)) for (i in 1:length(todrop))
if (length(todrop[[i]])>0) sl[[i]] = sl[[i]][,-todrop[[i]]] sms@smList = sl sms } library(GGdata)
library(multicore) data(eset) ex library(genefilter) exf1 = nsFilter(ex) length(get("17", revmap(illuminaHuman))
kp = intersect(ZZ, featureNames(exf1[[1]])) c17 = getSS("GGdata", "17", renameChrs="chr17", probesToKeep
f1dm = eqtlTests(c17, ~ male, targdir="c17c", genegran=1, geneApply=mclapply) save(f1dm, file="f1dm.rda")
set.seed(1234); permf1dm = eqtlTests(permEx(c17), ~ male, targdir="c17c_perm", genegran=1, geneApply=m
save(permf1dm, file="permf1dm.rda")

```

## Value

an instance of [eqtlTestsManager-class](#)

## Examples

```
f1 = observed17ceu()
f1
f1@call
f1[ rsid("rs7502145"), probeId("GI_10190685-S")]
pf1 = onePerm17ceu()
pf1[ rsid("rs7502145"), probeId("GI_10190685-S")]
```

rules.n43

*snpStats imputation rules instance*

## Description

snpStats imputation rules instance

## Usage

```
data(rules.n43)
```

## Format

The format is:

```
Formal class 'ImputationRules' [package "snpStats"] with 1 slots
..@ .Data:List of 470806
... ...$ : NULL
... ...$ :List of 4
... ...$ maf : num 0.128
... ...$ r.squared: num 0.901
... ...$ snps : chr [1:4] "rs11654695" "rs9789059" "rs8073513" "rs7225087"
... ...$ hap.probs: num [1:32] 0.00 1.02e-21 0.00 1.45e-07 1.52e-07 ...
... ...$ :List of 4
... ...$ maf : num 0.163
... ...$ r.squared: num 0.802
... ...$ snps : chr [1:4] "rs11654695" "rs12449775" "rs8078223" "rs9907102"
... ...$ hap.probs: num [1:32] 0.02863 0.0253 0.00342 0.07866 0.01717 ...
... ...$ :List of 4
... ...$ maf : num 0.116
... ...$ r.squared: num 0.881
... ...$ snps : chr [1:4] "rs11654695" "rs9789059" "rs8073513" "rs4968164"
... ...$ hap.probs: num [1:32] 0.00 1.96e-13 0.00 4.91e-05 1.05e-06 ...
... ...$ : NULL
... ... [list output truncated]
```

## Examples

```
library(snpStats)
data(rules.n43)
rules.n43[1:4]
## maybe str(rules.n43) ; plot(rules.n43) ...
```

---

snpgr17

*SNP locations for chr 17*

---

## Description

SNP locations for chr 17

## Usage

```
data(snpgr17)
```

## Format

The format is:

```
Formal class 'GRanges' [package "GenomicRanges"] with 6 slots
..@ seqnames :Formal class 'Rle' [package "IRanges"] with 4 slots
... ..@ values : Factor w/ 1 level "chr17": 1
... ..@ lengths : int 316396
... ..@ elementMetadata: NULL
... ..@ metadata : list()
..@ ranges :Formal class 'IRanges' [package "IRanges"] with 6 slots
... ..@ start : int [1:316396] 6934 7214 7242 8611 11743 11830 13546 13905 14122 14300 ...
... ..@ width : int [1:316396] 1 1 1 1 1 1 1 1 1 ...
... ..@ NAMES : chr [1:316396] "rs1106176" "rs6420494" "rs6420495" "rs62054996" ...
... ..@ elementType : chr "integer"
... ..@ elementMetadata: NULL
... ..@ metadata : list()
..@ strand :Formal class 'Rle' [package "IRanges"] with 4 slots
... ..@ values : Factor w/ 3 levels "+","-","*": 3
... ..@ lengths : int 316396
... ..@ elementMetadata: NULL
... ..@ metadata : list()
..@ elementMetadata:Formal class 'DataFrame' [package "IRanges"] with 6 slots
... ..@ rownames : NULL
... ..@ nrows : int 316396
... ..@ listData : Named list()
... ..@ elementType : chr "ANY"
... ..@ elementMetadata: NULL
... ..@ metadata : list()
..@ seqinfo :Formal class 'Seqinfo' [package "GenomicRanges"] with 3 slots
... ..@ seqnames : chr "chr17"
... ..@ seqlengths : int NA
... ..@ is_circular: logi NA
..@ metadata : list()
```

## Examples

```
data(snpgr17)
snpgr17
## maybe str(snpgr17) ; plot(snpgr17) ...
```

# Index

## \*Topic **datasets**

c17imp, [2](#)  
g17rngsnr, [3](#)  
rules.n43, [5](#)  
snpgr17, [6](#)

## \*Topic **models**

observed17ceu, [4](#)

## \*Topic **package**

ggtut-package, [1](#)

bestGenesPerSNP \_ obs (ggtut-package), [1](#)  
bestGenesPerSNP \_ perm (ggtut-package), [1](#)

c17imp, [2](#)

CPS17 (ggtut-package), [1](#)

g17rngsnr, [3](#)

getFixedBFL (ggtut-package), [1](#)

ggtut (ggtut-package), [1](#)

ggtut-package, [1](#)

hg18tx (ggtut-package), [1](#)

observed17ceu, [4](#)

onePerm17ceu (observed17ceu), [4](#)

PERMCPS17 (ggtut-package), [1](#)

permsb1 (ggtut-package), [1](#)

rsb2 \_ list (ggtut-package), [1](#)

rsb2 \_ perm \_ list (ggtut-package), [1](#)

rules.n43, [5](#)

sb1 (ggtut-package), [1](#)

snpgr17, [6](#)

SVA1 (ggtut-package), [1](#)

tr17 \_ 1 \_ 9 (observed17ceu), [4](#)

tr17 \_ 1 \_ 9 \_ perm (observed17ceu), [4](#)