

# Package ‘FunciSNP’

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**Type** Package

**Title** The Functional Integration of SNPs with Phenotype by Coincidence with Chromatin Biofeatures

**Version** 0.1.7

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**biocViews** Infrastructure, DataRepresentation, DataImport,SequenceMatching, Annotation

**Depends**

R (>= 2.14.0), Rsamtools (>= 1.6.1), rtracklayer(>= 1.14.1),GGtools (>= 4.0.0), methods, ChIP-peakAnno (>= 2.2.0),GenomicRanges, TxDb.Hsapiens.UCSC.hg19.knownGene,VariantAnnotation, plyr, org.Hs.eg.db, snp

**Imports** IRanges, AnnotationDbi

**Suggests** gplots (>= 2.10.1), ggplot2 (>= 0.8.9), matlab (>= 0.8.9)

**Enhances** parallel

**Description** FunciSNP integrates information from GWAS, 1000genomes and chromatin feature to identify functional SNP in coding or non-coding regions.

**License** GPL-3

**LazyLoad** yes

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FunciSNP-package	<i>Functional Identification of SNPs with Phenotype by Coincidence with Chromatin Biofeatures</i>
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**Description**

The package includes functions to identify and annotate putative functional SNPs using information derived from GWAS, 1000 genomes database, and sequences around peaks.

**Details**

Package:	FunciSNP
Type:	Package
Version:	0.1.7
Date:	2012-12-12
License:	GPL-3
LazyLoad:	yes

**Author(s)**

Simon Coetzee and Houtan Noushmehr  
Maintainer: Simon G. Coetzee <scoetzee@gmail.com>

**References**

Coetzee SG et al. submitted for review. 2012

**See Also**

[FunciSNPplot](#), [FunciSNPAnnotateSummary](#), [FunciSNPtable](#), [FunciSNPbed](#)

**Examples**

```
##  
## Glioblastoma analysis using FunciSNP  
##
```

```
## Full path to the example regions file for Glioblastoma
# (collected from SNPedia)
glioma.snp <- file.path(system.file('extdata',
  package='FunciSNP'),
  dir(system.file('extdata',package='FunciSNP'),
  pattern='.snp$'));

## Full path to the example biological features BED files
# derived from the ENCODE project for Glioblastoma U-87
# cell lines.
glioma.bio <- system.file('extdata',package='FunciSNP');

## FunciSNP analysis, extracts correlated SNPs from the
# 1000 genomes db ("ncbi") and finds overlaps between
# correlated SNP and biological features and then
# calculates LD (Rsquare, Dprime, distance, p-value).
# Do not run. Can take more than 5 min depending on internet connection and number of CPUs.
#glioma <- FunciSNP(snp.regions.file=glioma.snp,
# bio.features.loc = glioma.bio, bio.features.TSS=FALSE);

##
data(glioma);
class(glioma);
glioma;
summary(glioma);
```

---

CorrelatedSNPs-class    *Class "CorrelatedSNPs"*

---

## Description

Class for CorrelatedSNPs

## Objects from the Class

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

## Slots

chromosome: Object of class "integer" ~~  
 position: Object of class "integer" ~~  
 snpid: Object of class "character" ~~  
 ref.allele: Object of class "character" ~~  
 alt.allele: Object of class "character" ~~  
 overlapping.features: Object of class "GRanges" ~~  
 genotype: Object of class "CorrGeno" ~~

```

ALL.R.squared: Object of class "matrix" ~~
AFR.R.squared: Object of class "matrix" ~~
AMR.R.squared: Object of class "matrix" ~~
ASN.R.squared: Object of class "matrix" ~~
EUR.R.squared: Object of class "matrix" ~~
ALL.D.prime: Object of class "matrix" ~~
AFR.D.prime: Object of class "matrix" ~~
AMR.D.prime: Object of class "matrix" ~~
ASN.D.prime: Object of class "matrix" ~~
EUR.D.prime: Object of class "matrix" ~~
ALL.p.value: Object of class "list" ~~
AFR.p.value: Object of class "list" ~~
AMR.p.value: Object of class "list" ~~
ASN.p.value: Object of class "list" ~~
EUR.p.value: Object of class "list" ~~

```

## Methods

```

AFR.D.prime<- signature(x = "CorrelatedSNPs"): ...
AFR.D.prime signature(x = "CorrelatedSNPs"): ...
AFR.p.value<- signature(x = "CorrelatedSNPs"): ...
AFR.p.value signature(x = "CorrelatedSNPs"): ...
AFR.R.squared<- signature(x = "CorrelatedSNPs"): ...
AFR.R.squared signature(x = "CorrelatedSNPs"): ...
ALL.D.prime<- signature(x = "CorrelatedSNPs"): ...
ALL.D.prime signature(x = "CorrelatedSNPs"): ...
ALL.p.value<- signature(x = "CorrelatedSNPs"): ...
ALL.p.value signature(x = "CorrelatedSNPs"): ...
ALL.R.squared<- signature(x = "CorrelatedSNPs"): ...
ALL.R.squared signature(x = "CorrelatedSNPs"): ...
alt.allele<- signature(x = "CorrelatedSNPs"): ...
alt.allele signature(x = "CorrelatedSNPs"): ...
AMR.D.prime<- signature(x = "CorrelatedSNPs"): ...
AMR.D.prime signature(x = "CorrelatedSNPs"): ...
AMR.p.value<- signature(x = "CorrelatedSNPs"): ...
AMR.p.value signature(x = "CorrelatedSNPs"): ...
AMR.R.squared<- signature(x = "CorrelatedSNPs"): ...
AMR.R.squared signature(x = "CorrelatedSNPs"): ...

```

```

ASN.D.prime<- signature(x = "CorrelatedSNPs"): ...
ASN.D.prime signature(x = "CorrelatedSNPs"): ...
ASN.p.value<- signature(x = "CorrelatedSNPs"): ...
ASN.p.value signature(x = "CorrelatedSNPs"): ...
ASN.R.squared<- signature(x = "CorrelatedSNPs"): ...
ASN.R.squared signature(x = "CorrelatedSNPs"): ...
chr<- signature(x = "CorrelatedSNPs"): ...
chr signature(x = "CorrelatedSNPs"): ...
EUR.D.prime<- signature(x = "CorrelatedSNPs"): ...
EUR.D.prime signature(x = "CorrelatedSNPs"): ...
EUR.p.value<- signature(x = "CorrelatedSNPs"): ...
EUR.p.value signature(x = "CorrelatedSNPs"): ...
EUR.R.squared<- signature(x = "CorrelatedSNPs"): ...
EUR.R.squared signature(x = "CorrelatedSNPs"): ...
overlapping.features<- signature(x = "CorrelatedSNPs"): ...
overlapping.features signature(x = "CorrelatedSNPs"): ...
pop.genotype<- signature(x = "CorrelatedSNPs"): ...
pop.genotype signature(x = "CorrelatedSNPs"): ...
position<- signature(x = "CorrelatedSNPs"): ...
position signature(x = "CorrelatedSNPs"): ...
ref.allele<- signature(x = "CorrelatedSNPs"): ...
ref.allele signature(x = "CorrelatedSNPs"): ...
snpid<- signature(x = "CorrelatedSNPs"): ...
snpid signature(x = "CorrelatedSNPs"): ...

```

## Note

NA

## Author(s)

Simon Coetzee, Houtan Noushmehr

## References

Coetzee SG et al. submitted for review. 2012

## See Also

[FunciSNP](#), [FunciSNPplot](#), [FunciSNPAnnotateSummary](#), [FunciSNPtable](#), [FunciSNPbed](#)

## Examples

```
showClass("CorrelatedSNPs")
```

---

CorrGeno-class

*Class "CorrGeno"*

---

### Description

placeholder««<

### Objects from the Class

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

placeholder««<

### Slots

**Snpmatrix:** Object of class "SnpMatrix"

**populations:** Object of class "list" placeholder««<

placeholder««<

### Methods

No methods defined with class "CorrGeno" in the signature. placeholder««<

### Note

placeholder««<

### Author(s)

placeholder««<

### References

placeholder««<

### See Also

placeholder««<

### Examples

```
showClass("CorrGeno")
```

---

FunciSNPAnnotateSummary

*Genomic Annotation of Func-y-SNPs.*

---

## Description

This will annotate all identified Func-y-SNP for it's distance to the nearest known TSS, whether it overlaps a known exon, intron, 5'UTR, 3'UTR, promoter, lincRNA or in gene desert (intergenic) regions.

## Usage

```
FunciSNPAnnotateSummary(snp.list)
```

## Arguments

snp.list	a FunciSNP object: snp.list represents the FunciSNP object output from FunciSNP. See <a href="#">FuncySNP</a> .
----------	---

## Details

All known genomic features (exon, intron, 5'UTR, 3'UTR, promoter, lincRNA or in gene desert (intergenic)) are used to annotate the newly identified Func-y-SNP. Information described in this data.frame is used for all summary plots, table, and bed file generations.

## Value

data.frame with rows for each correlated SNP.

## Note

NA

## Author(s)

Simon Coetzee, Houtan Noushmehr

## References

Coetzee SG et al. submitted for review. 2012

## See Also

[FuncySNP](#), [FunciSNPplot](#), [FunciSNPAnnotateSummary](#), [FunciSNPtable](#), [FunciSNPbed](#)

## Examples

```
data(glioma);
gl <- FunciSNPAnnotateSummary(glioma);
dim(gl)
head(gl)
names(gl)
```

---

FunciSNPbed	<i>Creates a BED file to view Func-y-SNPs in your favorite genome browser</i>
-------------	---

---

## Description

FunciSNPbed will output a BED file to a specified folder. The BED file is in standard UCSC Genome Browser format (<http://genome.ucsc.edu/FAQ/FAQformat>). Each tagSNP is colored black and each Func-y-SNP is colored red.

## Usage

```
FunciSNPbed(dat, rsq, path = getwd(), filename = NULL)
```

## Arguments

dat	FunciSNP data.frame: dat is a data.frame object from FunciSNPAnnotateSummary. Need to run <a href="#">FunciSNPAnnotateSummary</a> first.
rsq	an interger (0-1): rsq is the Rsquared cutoff used to subset.
path	a character: path is the path to the folder where to save the BED file. Default to getwd() or current working directory.
filename	a character: filename is the name of the BED file. If NULL, filename is 'FunciSNP_results_rsq.RSQ value.bed'

## Details

FunciSNPbed outputs a unique BED file which can be used to view in any genomic browser compatible with BED formats. To learn more about BED formats, see UCSC Genome Browser FAQ (<http://genome.ucsc.edu/FAQ/FAQformat>). Each tagSNP which is in LD to a corresponding Func-y-SNP overlapping at least one biofeature is colored black, while the Func-y-SNP is colored red. The initial position is provided by the first tagSNP and the first linked Func-y-SNP. We recommend using UCSC genome browser to view your BED files. This is useful so you can view all public and private tracks in relation to FunciSNP results.

## Value

BED file is outputted as a tab-delimited file in the specified 'path' folder. See example below.

## Note

NA



**Author(s)**

Simon Coetzee, Houtan Noushmehr

**References**

Coetzee SG et al. submitted for review. 2012

**See Also**

[FuncySNP](#), [FunciSNPplot](#), [FunciSNPAnnotateSummary](#), [FunciSNPtable](#), [FunciSNPbed](#)

**Examples**

```
##
data(glioma);
glioma.anno <- FunciSNPAnnotateSummary(glioma);
FunciSNPbed(glioma.anno, rsq=0.9);
####
#Bed file "FunciSNP_results_rsqu.0.9.bed" created successfully.
#(See folder: "/home/houtan/Downloads/")
#Total corSNP (RED): 15
#Total tagSNP (BLK): 1

#To view results, submit bed file as a
# custom track in UCSC Genome Browser (genome.ucsc.edu),

#Now have fun with your new Func-y SNPs!!
####
```

---

FunciSNPidsFromSummary

*coming soon.*

---

**Description**

placeholder««<

**Usage**

```
FunciSNPidsFromSummary(dat, tagsnpid = NULL, num.features, rsq = 0)
```

**Arguments**

dat	placeholder««<
tagsnpid	placeholder««<
num.features	placeholder««<
rsq	placeholder««< placeholder««<

Details

placeholder««<

Value

placeholder««<

Note

NA

Author(s)

Simon Coetzee, Houtan Noushmehr

References

Coetzee SG et al. submitted for review. 2012

See Also

[FunciSNP](#), [FunciSNPplot](#), [FunciSNPAnnotateSummary](#), [FunciSNPtable](#), [FunciSNPbed](#)

Examples

## coming soon

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FunciSNPplot	<i>FunciSNPplot to visualize Func-y-SNP summary.</i>
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---

Description

FunciSNPplot is a function developed to plot various types of plots to summarize and assist end-user in making informed discoveries of FunciSNP results. Plots can be stored in a folder for future reference.

Usage

FunciSNPplot(dat, rsq = 0, split = FALSE, splitbysnp = FALSE, tagSummary = FALSE, heatmap = FALSE, genom

**Arguments**

<code>dat</code>	FunciSNP data.frame: <code>dat</code> is a data.frame object from <code>FunciSNPAnnotateSummary</code> . Need to run <a href="#">FunciSNPAnnotateSummary</a> first.
<code>rsq</code>	an interger (0-1): <code>rsq</code> is the Rsquared cutoff used to subset.
<code>split</code>	logical: <code>split</code> will generate distribution plot of all Correlated SNPs by Rsquare values.
<code>splitbysnp</code>	logical: <code>splitbysnp</code> is similar to <code>split</code> but instead split the distribution by tagSNP.
<code>tagSummary</code>	logical: <code>tagSummary</code> Will output two plots per biofeature. The first one is a scatter plot showing the relationship between Rsquare and Distance to tagSNP for each Func-y-SNP. The second plot is a histogram distribution of number of correlated SNPs at each Rsquare value. Each set of plot is further divided by tagSNP. Best if used with <code>rsq</code> value.
<code>heatmap</code>	logical: <code>heatmap</code> correlation heatmap to visualize the number of correlated SNPs at each tagSNP overlapping each biological feature. Most informative if used with a <code>rsq</code> value.
<code>genomicSum</code>	logical: <code>genomicSum</code> Stacked bar chart summarizing all correlated SNPs for each of the identified genomie features (exon, intron, 5'UTR, 3'UTR, promoter, lincRNA or in gene desert (intergenic)). Most informative if used with a <code>rsq</code> value.
<code>save</code>	logical: <code>save</code> to save outputs to folder. Set at <code>getwd()</code> , in folder 'FunciSNP.VERSION/plots
<code>pathplot</code>	a character: <code>pathplot</code> is the path to the folder where to save the plots. Default to <code>getwd()</code> or current working directory.

**Details**

NA

**Value**

Plots are generated either in X11 or in specified folder.

**Note**

NA

**Author(s)**

Simon Coetzee, Houtan Noushmehr

**References**

Coetzee SG et al. submitted for review. 2012

**See Also**[FunciSNP](#), [FunciSNPplot](#), [FunciSNPAnnotateSummary](#), [FunciSNPtable](#), [FunciSNPbed](#)

**Examples**

```

data(glioma)
gl <- FunciSNPAnnotateSummary(glioma)
FunciSNPplot(gl)
FunciSNPplot(gl, rsq=0, genomicSum=TRUE, save=FALSE)
FunciSNPplot(gl, rsq=0.5, genomicSum=TRUE, save=FALSE)
# DO NOT RUN
#FunciSNPplot(gl, tagSummary=TRUE, rsq=0.5)
#

```

---

FunciSNPSummaryOverlaps

*coming*

---

**Description**

placeholder««<

**Usage**

```
FunciSNPSummaryOverlaps(dat, rsq = 0)
```

**Arguments**

dat	placeholder««<
rsq	placeholder««< placeholder««<

**Details**

placeholder««<

**Value**

placeholder««<

**Note**

NA

**Author(s)**

Simon Coetzee, Houtan Noushmehr

**References**

Coetzee SG et al. submitted for review. 2012

**See Also**

[FunciSNP](#), [FunciSNPplot](#), [FunciSNPAnnotateSummary](#), [FunciSNPtable](#), [FunciSNPbed](#)

**Examples**

```
##coming soon.
```

---

FunciSNPtable	<i>Will output a summary report from FunciSNP at specified Rsquare cutoffs.</i>
---------------	---

---

**Description**

Using a specified Rsquare value (0-1) to subset the data, a table is generated which summarizes the total number of Func-y-SNPs, associated tagSNPs, and number of overlapping biofeatures.

**Usage**

```
FunciSNPtable(dat, rsq, geneSum = FALSE)
```

**Arguments**

dat	FunciSNP data.frame: dat is a data.frame object from FunciSNPAnnotateSummary. Need to run <a href="#">FunciSNPAnnotateSummary</a> first.
rsq	an interger (0-1): rsq is the Rsquared cutoff used to subset.
geneSum	logical: geneSum is set to FALSE. Setting to TRUE will output a list of Gene names which are nearest to the Func-y-SNP.

**Details**

Using a specified Rsquare value (0-1) to subset the data, a table is generated which summarizes the total number of Func-y-SNPs, associated tagSNPs, and number of overlapping biofeatures. This will provide user a first look at the total number of available Func-y-SNP at a particular Rsquare cutoff. If geneSum is set to TRUE, a list of gene names is reported instead.

**Value**

Standard output which summarizes FunciSNP results.

**Note**

NA

**Author(s)**

Simon Coetzee, Houtan Noushmehr

## References

Coetzee SG et al. submitted for review. 2012

## See Also

[FuncySNP](#), [FunciSNPplot](#), [FunciSNPAnnotateSummary](#), [FunciSNPtable](#), [FunciSNPbed](#)

## Examples

```
data(glioma);
gl <- FunciSNPAnnotateSummary(glioma);
FunciSNPtable(gl, rsq=0.5);
FunciSNPtable(gl, rsq=0.5, geneSum=TRUE);
```

---

FuncySNP

*Functional Identification of SNPs with Phenotype by Coincidence with Chromatin Biofeatures*

---

## Description

Given a set of known tag-SNPs associated with a particular phenotype (e.g. disease, trait), and a set of available biological features (e.g. peaks derived from ChIP-seq experiments for phenotype), returns correlated SNPs (from the 1000 genomes db) which are in linkage disequilibrium (LD) to a known disease associated tag-SNP and overlaps chromatin biological features. These identified correlated SNPs are characterized as putative functional SNPs for a particular trait.

## Usage

```
FuncySNP(snp.regions.file, bio.features.loc = NULL,
         bio.features.TSS = TRUE,
         par.threads=detectCores()/2,
         verbose = par.threads < 2, method.p = "BH",
         reduce.by.features = TRUE, search.window = 200000)
```

## Arguments

snp.regions.file

path: Location of the regions file: Regions file is tab-delimited and contains three elements per row. First element defines the genomic location of the tagSNP, 'chr:position' (e.g. 5:5420030). Second element contains the tagSNP name, 'rsID' (e.g. rs6010620). Third element defines the 'POPULATION' (ASN, EUR, AFR, ALL) where the tagSNP was identified (e.g. ASN, EUR, AFR, ALL).

SNP Region file is imported and each row element (tagSNP element) is parsed for tagSNP name (rsXXXX), population (ASN, EUR, AFR, or ALL), and genomic location. Genomic location is used to define the window size (see 'search.window')

	argument). See example file here: <code>file.path(system.file('data',package='FunciSNP'), dir(system.file('data',package='FunciSNP'), pattern='.snp\$'))</code> ;
<code>bio.features.loc</code>	path: Location of the biological features folder: Each biological feature for a particular genomic phenotype should be separated as individual BED files (tab delimited file with chr, start and end). See UCSC for more information about BED formats <a href="http://genome.ucsc.edu/FAQ/FAQformat.html#format1">http://genome.ucsc.edu/FAQ/FAQformat.html#format1</a> . See example below. Default set to NULL.
<code>bio.features.TSS</code>	logical: To include promoter regions as an additional biofeature in the analysis. Promoters defined as -1000 to +100 bp of a known TSS. File extracted on Feb. 9, 2012 from UCSC genome table browser. Default set to TRUE.
<code>par.threads</code>	an integer: Number of CPU cores to use for FunciSNP analysis. Default set at <code>detectCores()/2</code> . If <code>par.threads &gt; 1</code> , then by default "verbose" = FALSE.
<code>verbose</code>	logical: If set to TRUE, then regardless of <code>par.threads</code> value, all verbose message will output to terminal. If set to FALSE, no verbose message will output to terminal, except for warnings(). Default setting depends on number of 'par.threads' value.
<code>method.p</code>	method: p-value correction (or adjustment) method (see <code>?p.adjust</code> ). Default set at "BH" (Benjamini & Hochberg (1995)).
<code>reduce.by.features</code>	logical: If set to TRUE, then only correlated SNPs overlapping biological features will be filtered and used to calculate Rsquared, Dprime, distance and p-value. In addition, only these correlated SNPs will be used to generate plots and summary analysis. If set to FALSE, all correlated SNPs regardless of overlap with biological features will have an associated Rsquare, Dprime, distance and p-value associated with the tag-SNP as defined by the 'search.window'.
<code>search.window</code>	an integer: genomic window size used to extract all available correlated SNPs from the 1000 genomes db. The window size is centered around the tagSNP position as defined in the <code>regions.file</code> .

## Details

This is the main function of FunciSNP. It will identify correlated SNPs which are in linkage disequilibrium (LD) to a known disease associated tagSNP. It will also determine if the correlated SNP in LD to the tagSNP overlaps a genomic biological feature. Correlated SNPs are directly imported from the current public release of the 1000 genomes database. 1000 genomes ftp servers available for the 1000 genomes public data: 1) National Center for Biotechnology Information (NCBI) <ftp://ftp-trace.ncbi.nih.gov/1000genomes/>; 2) European Bioinformatics Institute (EBI) <ftp://ftp.1000genomes.ebi.ac.uk/vol1/>.

Correlated SNPs in LD to a tagSNP and overlapping genomic biological features are known as putative functional SNPs (also defined as 'Func-y-SNP' elsewhere in the package.).

## Value

TSList	FunciSNP object.
--------	------------------

**Note**

NA

**Author(s)**

Simon Coetzee, Houtan Noushmehr

**References**

Coetzee SG et al. submitted for review. 2012

**See Also**[FunciSNPplot](#), [FunciSNPAnnotateSummary](#), [FunciSNPtable](#), [FunciSNPbed](#)**Examples**

```
##
## Glioblastoma analysis using FunciSNP
##
## Full path to the example regions file for Glioblastoma
# (collected from SNPedia)
glioma.snp <- file.path(system.file('extdata',
  package='FunciSNP'),
  dir(system.file('extdata',package='FunciSNP'),
  pattern='.snp$'));

## Full path to the example biological features BED files
# derived from the ENCODE project for Glioblastoma U-87
# cell lines.
glioma.bio <- system.file('extdata',package='FunciSNP');

## FunciSNP analysis, extracts correlated SNPs from the
# 1000 genomes db ("ncbi") and finds overlaps between
# correlated SNP and biological features and then
# calculates LD (Rsquare, Dprime, distance, p-value).
# Do not run. Can take more than 5 min depending on internet connection and number of CPUs.
#glioma <- FuncySNP(snp.regions.file=glioma.snp,
# bio.features.loc = glioma.bio, bio.features.TSS=FALSE);

##
data(glioma);
class(glioma);
glioma;
summary(glioma);
```



TagSNP-class

Class "TagSNP"

**Description**

hello

**Objects from the Class**

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

**Slots**

chromosome: Object of class "integer" ~~  
 position: Object of class "integer" ~~  
 snpid: Object of class "character" ~~  
 population: Object of class "character" ~~  
 ref.allele: Object of class "character" ~~  
 alt.allele: Object of class "character" ~~  
 overlapping.features: Object of class "GRanges" ~~  
 genotype: Object of class "SnpMatrix" ~~  
 R.squared.corrsnps: Object of class "dgCMatrix" ~~  
 D.prime.corrsnps: Object of class "dgCMatrix" ~~  
 correlated.snps: Object of class "CorrelatedSNPs" ~~

**Methods**

**AFR.overlapping.snps.geno** signature(object = "TagSNP"): ...  
**ALL.overlapping.snps.geno** signature(object = "TagSNP"): ...  
**alt.allele<-** signature(x = "TagSNP"): ...  
**alt.allele** signature(x = "TagSNP"): ...  
**AMR.overlapping.snps.geno** signature(object = "TagSNP"): ...  
**ASN.overlapping.snps.geno** signature(object = "TagSNP"): ...  
**chr<-** signature(x = "TagSNP"): ...  
**chr** signature(x = "TagSNP"): ...  
**correlated.snps<-** signature(x = "TagSNP"): ...  
**correlated.snps** signature(x = "TagSNP"): ...  
**D.prime.corrsnps<-** signature(x = "TagSNP"): ...  
**D.prime.corrsnps** signature(x = "TagSNP"): ...  
**EUR.overlapping.snps.geno** signature(object = "TagSNP"): ...

```
genotype<- signature(x = "TagSNP"): ...  
genotype signature(x = "TagSNP"): ...  
overlapping.features<- signature(x = "TagSNP"): ...  
overlapping.features signature(x = "TagSNP"): ...  
population<- signature(x = "TagSNP"): ...  
population signature(x = "TagSNP"): ...  
position<- signature(x = "TagSNP"): ...  
position signature(x = "TagSNP"): ...  
ref.allele<- signature(x = "TagSNP"): ...  
ref.allele signature(x = "TagSNP"): ...  
R.squared.corrsnps<- signature(x = "TagSNP"): ...  
R.squared.corrsnps signature(x = "TagSNP"): ...  
show signature(object = "TagSNP"): ...  
snpid<- signature(x = "TagSNP"): ...  
snpid signature(x = "TagSNP"): ...
```

#### Note

NA

#### Author(s)

Simon Coetzee, Houtan Noushmehr

#### References

Coetzee SG et al. submitted for review. 2012

#### See Also

[FancySNP](#), [FunciSNPplot](#), [FunciSNPAnnotateSummary](#), [FunciSNPtable](#), [FunciSNPbed](#)

#### Examples

```
showClass("TagSNP")
```

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TSList-class	Class "TSList"
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**Description**

ffff

**Objects from the Class**

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

**Slots**

snp.data: Object of class "list" ~~  
summary.data: Object of class "data.frame" ~~  
elementType: Object of class "character" ~~  
elementMetadata: Object of class "DataTableORNULL" ~~  
metadata: Object of class "list" ~~

**Methods**

**show** signature(object = "TSList"): ...  
**summary** signature(object = "TSList"): ...

**Note**

NA

**Author(s)**

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**References**

Coetzee SG et al. submitted for review. 2012

**See Also**

[FuncySNP](#), [FunciSNPplot](#), [FunciSNPAnnotateSummary](#), [FunciSNPtable](#), [FunciSNPbed](#)

**Examples**

```
showClass("TSList")
```

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