Package 'myFun'

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Type Package

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Description My utility functions for R	
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adjustPositions

adjustPositions

Description

Adjust genomic positions

Usage

```
adjustPositions(
   DF,
   CHRsize,
   chr_column = "chr",
   start_column = "start",
   end_column = "end",
   suffix = "_adj"
)
```

Arguments

```
DF a data.frame

CHRsize a data.frame from the load_CHRsize function

chr_column a column name with chromosome information (default: "chr")

start_column a column name with start position (default: "start")

end_column a column name with end position (default: "end")

suffix a suffix for the adjusted positions (default: "_adj")
```

Details

This function adjusts genomic positions according to the chromosome sizes. The first nucleotide of chromosome 2 corresponds to the size of the chromosome 1 + 1bp and so on.

Value

A data.frame with adjusted genomic positions

Author(s)

tlesluyes

Examples

```
DF=data.frame(chr=c(1:3), start=rep(1e6, 3), end=rep(125e6, 3))
load_CHRsize("hg19")
adjustPositions(DF, CHRsize)
```

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checkGRlist

checkGRlist

Description

Check that the given object is a list of GRanges objects

Usage

```
checkGRlist(myGRList)
```

Arguments

myGRList

a list of GRanges objects

Details

This function checks that the given object is a list of GRanges objects.

Value

TRUE if the input is a list of GRanges objects

Author(s)

tlesluyes

Examples

```
require(GenomicRanges)
GR1=GRanges(seqnames="1", ranges=IRanges(start=1, end=1000))
GR2=GRanges(seqnames="1", ranges=IRanges(start=10, end=2000))
checkGRlist(list(GR1, GR2))
```

 ${\tt computeISA}$

computeISA

Description

Compute the inter-sample agreement (ISA)

Usage

```
computeISA(GR1, GR2, CNstatus = "CNstatus")
```

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Arguments

GR1 a GRanges object corresponding to a single CNA profile
GR2 a GRanges object corresponding to a single CNA profile

CNstatus a metadata column name for the copy-number status (default: "CNstatus"). Can

be total (e.g. "3") or allele-specific (e.g. "2+1")

Details

This function computes the inter-sample agreement (ISA) between two profiles (as GRanges objects). This corresponds to the fraction of the genome (%) with the same CN status.

Value

A percentage representing the ISA

Author(s)

tlesluyes

Examples

computeMD

computeMD

Description

Compute the Manhattan distance (MD)

Usage

```
computeMD(GR1, GR2, nMajor = "nMajor", nMinor = "nMinor", convertMb = FALSE)
```

Arguments

```
GR1 a GRanges object corresponding to a single CNA profile

GR2 a GRanges object corresponding to a single CNA profile

nMajor a metadata column name for the major allele (default: "nMajor")

nMinor a metadata column name for the minor allele (default: "nMinor")

convertMb a boolean, the MD will be converted to megabases if set to TRUE (default: FALSE)
```

Details

This function computes the Manhattan distance (MD) between two profiles (as GRanges objects).

Value

A numeric value representing the MD

Author(s)

tlesluyes

Examples

Description

Generate cytoband and CHRsize information

Usage

```
generate_cytoband_and_CHRsize(cytoband_file)
```

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Arguments

```
cytoband_file a cytoband file
```

Details

This function generates cytoband and CHRsize information from a cytoband file. This can be obtained from the UCSC table browser -> select a genome/assembly -> "Mapping and Sequencing" -> "Chromosome Band" (not the ideogram version!) -> "get output" -> Remove the first "#" character (keep the header!).

Value

A list with both the cytoband and CHRsize information

Author(s)

tlesluyes

See Also

```
load_CHRsize("hg38"); load_cytoband("hg38")
```

harmonizeGRanges

harmonizeGRanges

Description

Harmonize GRanges objects

Usage

harmonizeGRanges(myGRList)

Arguments

myGRList

a list of GRanges objects, each object should correspond to one CNA profile

Details

This function harmonizes GRanges objects by keeping only regions covered by all samples.

Value

A list of harmonized GRanges objects

Author(s)

tlesluyes

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Examples

```
require(GenomicRanges)
GR1=GRanges(seqnames="1", ranges=IRanges(start=1, end=1000), nMajor=1, nMinor=1)
GR2=GRanges(seqnames="1", ranges=IRanges(start=10, end=2000), nMajor=2, nMinor=1)
harmonizeGRanges(list(GR1, GR2))
```

load_CHRsize

load_CHRsize

Description

Load CHRsize information

Usage

```
load_CHRsize(assembly)
```

Arguments

assembly

an assembly (hg19 or hg38)

Details

This function loads CHRsize information for a given assembly. It is then available as a data.frame called CHRsize in the environment.

Value

A data.frame with the CHRsize information

Author(s)

tlesluyes

Examples

```
load_CHRsize("hg38"); head(CHRsize)
```

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 $load_cytoband$

load_cytoband

Description

Load cytoband information

Usage

```
load_cytoband(assembly)
```

Arguments

assembly

an assembly (hg19 or hg38)

Details

This function loads cytoband information for a given assembly. It is then available as a data.frame called cytoband in the environment.

Value

A data.frame with the cytoband information

Author(s)

tlesluyes

Examples

```
load_cytoband("hg38"); head(cytoband)
```

occurrenceGRanges

occurrenceGRanges

Description

Get the occurrence of events

Usage

```
occurrenceGRanges(myGRList, myMetadata)
```

Arguments

myGRList a list of GRanges objects, each object should correspond to one CNA profile

myMetadata a vector of metadata to consider

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Details

This function gets the occurrence of events in a list of GRanges objects. All objects must have the same metadata columns and metadata must be TRUE/FALSE.

Value

A GRanges object with nSamples as the total number of samples and metadata columns with the occurrence of events

Author(s)

tlesluyes

Examples

```
require(GenomicRanges)
GR1=GRanges(seqnames="1", ranges=IRanges(start=1, end=1000), Gain=TRUE, Loss=FALSE)
GR2=GRanges(seqnames="1", ranges=IRanges(start=10, end=2000), Gain=FALSE, Loss=TRUE)
occurrenceGRanges(list(GR1, GR2),c("Gain", "Loss"))
```

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