# Package 'myFun'

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Type Package
Title myFun is a collection of my favorite R functions, packaged for simplicity
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<b>Description</b> My utility functions for R.
<pre>URL https://github.com/tlesluyes/myFun</pre>
<pre>BugReports https://github.com/tlesluyes/myFun/issues</pre>
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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.

checkGRlist 3

### 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)
```

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

```
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))
```

4 computeBAF

### Description

Compute the theoretical BAF values for a given segment

### Usage

```
computeBAF(nMajor, nMinor, purity, digits = 4)
```

### **Arguments**

nMajor	the number of copies of the major allele
nMinor	the number of copies of the minor allele
purity	the purity estimate of the tumour
digits	a numeric, the number of digits to round to (default: 4)

### **Details**

This function computes the theoretical BAF values for a given segment (from nMajor, nMinor and purity values).

#### Value

A vector of two numbers representing the BAF values

### Author(s)

tlesluyes

#### See Also

https://doi.org/10.1038/s41592-020-01013-2

```
# A 2+1 state in a tumour with 90% purity computeBAF(2, 1, 0.9) # A 1+0 state in a tumour with 60% purity computeBAF(1, 0, 0.6)
```

computeFit 5

### Description

Compute the purity/ploidy fit for a given segment

#### Usage

```
computeFit(logR, BAF, nMajor, nMinor, gamma, digits = 4)
```

### Arguments

logR	the logR value of the segment
BAF	the BAF value of the segment (upper band only so the value should be in the $0.5\text{-}1\ \text{space})$
nMajor	the number of copies of the major allele
nMinor	the number of copies of the minor allele
gamma	the gamma parameter is platform-dependent and represents the expected logR decrease in a diploid sample where one copy is lost (should be 1 for HTS data and 0.55 for SNP arrays)
digits	a numeric, the number of digits to round to (default: 4)

#### **Details**

This function computes the purity/ploidy fit (rho, psi and psit) for a given segment (from logR, BAF, proposed nMajor and proposed nMinor).

#### Value

A list with the rho (=purity), psi (=total ploidy) and psit (=tumour ploidy) values

### Author(s)

tlesluyes

### See Also

https://doi.org/10.1038/s41592-020-01013-2

```
# A segment has logR=0.5361 and BAF=0.3448/0.6552 # What is the purity/ploidy fit if I believe that the segment is 2+1? computeFit(0.5361, 0.6552, 2, 1, 1) # purity=90%; ploidy=2
```

6 computeISA

computeISA

### Description

Compute the inter-sample agreement (ISA)

#### Usage

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

#### **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

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computeISA_batch	computeISA_	batch
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#### **Description**

Compute the inter-sample agreement (ISA) for a batch of samples

### Usage

```
computeISA_batch(myGRList, cores = 1, min_seg_size = 0, CNstatus = "CNstatus")
```

### **Arguments**

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

cores a numeric, the number of cores to use (default: 1)

min\_seg\_size a numeric, the minimum segment size (in bp) to consider (default: 0)

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 multiple profiles (as a list of GRanges objects).

#### Value

A matrix of ISA values

#### Author(s)

tlesluyes

8 computeLogR

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### **Description**

Compute the theoretical logR value for a given segment

### Usage

```
computeLogR(nMajor, nMinor, purity, ploidy, digits = 4)
```

### **Arguments**

nMajor	the number of copies of the major allele
nMinor	the number of copies of the minor allele
purity	the purity estimate of the tumour
ploidy	the ploidy estimate of the tumour
digits	a numeric, the number of digits to round to (default: 4)

#### **Details**

This function computes the theoretical logR value for a given segment (from nMajor, nMinor, purity and ploidy values). Since logR isn't allele-specific, ntot can be used instead of nMajor (and nMinor should set to 0).

#### Value

A number representing the logR value

### Author(s)

tlesluyes

### See Also

https://doi.org/10.1038/s41592-020-01013-2

```
# A 2+1 state in a diploid tumour with 90% purity
computeLogR(2, 1, 0.9, 2)
# A loss of 1 copy (2+1) in a pseudo-tetraploid tumour with 60% purity
computeLogR(2, 1, 0.6, 3.5)
```

computeMD 9

### **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

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computeMD\_batch

computeMD\_batch

### **Description**

Compute the Manhattan distance (MD) for a batch of samples

### Usage

```
computeMD_batch(
  myGRList,
  cores = 1,
  min_seg_size = 0,
  nMajor = "nMajor",
  nMinor = "nMinor",
  convertMb = FALSE
)
```

#### **Arguments**

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

cores a numeric, the number of cores to use (default: 1)

min\_seg\_size a numeric, the minimum segment size (in bp) to consider (default: 0)

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 multiple profiles (as a list of GRanges objects).

#### Value

A matrix of MD values

### Author(s)

tlesluyes

#### **Examples**

```
require("GenomicRanges")
GR1=GRanges(segnames=rep("1", 3),
            ranges=IRanges(start=c(1, 1001, 10001), end=c(1000, 10000, 20000)),
            nMajor=c(1, 2, 1),
            nMinor=c(1, 1, 1))
GR2=GRanges(seqnames=rep("1", 2),
            ranges=IRanges(start=c(500, 10001), end=c(10000, 25000)),
            nMajor=c(2, 1),
            nMinor=c(1, 1)
GR3=GRanges(seqnames="1",
            ranges=IRanges(start=500, end=25000),
            nMajor=1,
            nMinor=1)
myGRList=list(GR1, GR2, GR3)
names(myGRList)=c("GR1", "GR2", "GR3")
computeMD_batch(myGRList)
```

### Description

Generate cytoband and CHRsize information

#### Usage

```
generate_cytoband_and_CHRsize(cytoband_file)
```

### **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

get\_all\_paths

#### See Also

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

```
get_all_paths get_all_paths
```

#### **Description**

Get all possibles paths between two copy-number states

#### Usage

```
get_all_paths(start, end, WGD, max_path_size = 5, simplify = TRUE)
```

### Arguments

start	a vector of length 2 (representing a copy-number state; e.g. $c(1, 1)$ represents a $1+1$ state), defining where to start
end	a vector of length 2 (representing a copy-number state; e.g. $c(1, 1)$ represents a $1+1$ state), defining where to end
WGD	a boolean defining if WGD events are allowed
max_path_size	an integer defining the maximum path size
simplify	a boolean defining if consecutive and opposite alterations (e.g. $+1/+0$ and then $-1/-0$ ) are allowed

### **Details**

This function returns all possible paths between two copy-number states. The expected input is allele-specific (with two values), but it can be used for total copy-number by setting c(ntot, 0). Possible events include: +1/+0 (gain of the major allele), -1/-0 (loss of the major allele), +0/+1 (gain of the minor allele), -0/-1 (loss of the minor allele) and WGD.

### Value

A vector of all possible paths given as characters (separator=";")

### Author(s)

tlesluyes

```
# Diploid baseline (1+1) turns into 2+1
print(get_all_paths(start=c(1, 1), end=c(2, 1), WGD=TRUE))
# Chromosome X in males (1+0) is gained (5 copies)
print(get_all_paths(start=c(1, 0), end=c(5, 0), WGD=TRUE))
```

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<pre>get_shortest_path</pre>
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### **Description**

Get the shortest path among several

### Usage

```
get_shortest_path(paths, wanted_WGD = NA, count_WGD = FALSE)
```

#### **Arguments**

paths all possible paths to consider

wanted\_WGD a numeric value defining the number of WGD events wanted (can be NA to

allow for any possibility, including no event at all; default: NA)

count\_WGD a boolean defining if the number of WGD events should be counted (default:

FALSE)

#### **Details**

This function returns the shortest possible path. It should be used after running the get\_all\_paths function or can be used as long as the input format is correct.

#### Value

A numeric value representing the minimal number of events, its name represents the full path

#### Author(s)

tlesluyes

```
# Diploid baseline (1+1) turns into 2+1
print(get_shortest_path(get_all_paths(start=c(1, 1), end=c(2, 1), WGD=TRUE)))
# Chromosome X in males (1+0) is gained (5 copies)
print(get_shortest_path(get_all_paths(start=c(1, 0), end=c(5, 0), WGD=TRUE)))
```

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harmonizeGRanges

harmonizeGRanges

### **Description**

Harmonize GRanges objects

### Usage

```
harmonizeGRanges(myGRList, cores = 1)
```

#### **Arguments**

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

cores a numeric, the number of cores to use (default: 1)

### **Details**

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

#### Value

A list of harmonized GRanges objects

### Author(s)

tlesluyes

### **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)
```

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### **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)
```

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

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#### **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

### **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

```
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"))
```

reestimate\_ploidy 17

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### Description

Compute the re-estimated ploidy for a given sample

### Usage

```
reestimate_ploidy(rho.old, psit.old, rho.new, WGD, digits = 4)
```

### Arguments

```
rho.old old purity estimate

psit.old old ploidy estimate

rho.new new purity estimate

WGD number of WGD events (0 if there is no WGD)

digits a numeric, the number of digits to round to (default: 4)
```

#### **Details**

This function computes the re-estimated ploidy for a given sample (from its old purity/ploidy fit and the re-estimated purity).

### Value

A number representing the re-estimated ploidy

### Author(s)

tlesluyes

```
# A pseudo-diploid sample has purity=74% and ploidy=2.4
# What is the re-estimated ploidy if I believe that the sample has purity=61%?
reestimate_ploidy(0.74, 2.4, 0.61, 0)
```

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reestimate_purity	reestimate_	purity
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### Description

Compute the re-estimated purity for a given sample

#### Usage

```
reestimate_purity(rho.old, psit.old, switch, digits = 4)
```

### Arguments

rho.old old purity estimate

psit.old old ploidy estimate

switch a character ("double" or "halve") indicating whether the ploidy should be doubled or halved

digits a numeric, the number of digits to round to (default: 4)

#### **Details**

This function computes the re-estimated purity for a given sample in the context of a jump in ploidy (so the matched ploidy needs to be doubled or halved).

### Value

A number representing the re-estimated purity

### Author(s)

tlesluyes

```
# A sample has purity=74% and ploidy=2.4 but the CNA profile needs to be doubled # What is the re-estimated purity? reestimate_purity(0.74, 2.4, "double")
```

splitDF

### Description

Split a data.frame

### Usage

```
splitDF(DF, chunks, shuffle = FALSE, seed = 1234)
```

### **Arguments**

DF a data.frame to split

chunks a number of chunks to obtain

shuffle a boolean, whether to shuffle the data.frame before splitting (default: FALSE)

seed a number, the seed for the random number generator (default: 1234)

### **Details**

This function splits a data.frame into a list of data.frames.

### Value

A list of data.frames

### Author(s)

tlesluyes

```
DF=data.frame(a=1:26, b=letters)
splitDF(DF, 3)
```

 ${\tt summarise\_segmetation} \ \ \textit{summarise\_segmetation}$ 

### **Description**

Summarise segmentation data

### Usage

```
summarise_segmetation(DF, col_chr, col_start, col_end, col_values)
```

### Arguments

DF	a data.frame with segmentation data
col_chr	a string, the name of the column containing the chromosome
col_start	a string, the name of the column containing the start position
col_end	a string, the name of the column containing the end position (can be the same as col_start for SNP-based segmentation where start=end)
col_values	a vector of strings, the names of the columns containing the values of interest $(logR,BAF,etc.)$

#### **Details**

This function summarises segmentation data, typically logR and/or BAF values for individual SNPs or loci.

#### Value

A named list with segments being a data.frame with the summarised information and IDs being a list of SNPs/loci associated with the different segments

### Author(s)

tlesluyes

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