

Package ‘tugHall.3’

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Title R-based script to simulate the cancer cell evolution

Version 3.0

Description tugHall (tumor gene-Hallmark) is a cancer-cell evolution model simulator, wherein gene mutations are linked to the hallmarks of cancer, which influence tumor cell behaviors.

License GPL (>= 3)

Depends R (>= 3.6.0)

Imports actuar,
graphics,
grDevices,
methods,
randomcoloR,
stats,
stringr,
utils

Suggests rmarkdown,
knitr,
testthat

Encoding UTF-8

Roxygen list(markdown = TRUE)

RoxygenNote 7.1.2

LazyData true

VignetteBuilder knitr

R topics documented:

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<i>add_deletion</i>	<i>Function to add deletion to gene map (chromosomal location data frame)</i>
---------------------	---

Description

Function to add deletion to gene map (chromosomal location data frame)

Usage

```
add_deletion(gm, Ref_start, Ref_end, Chr)
```

Arguments

<code>gm</code>	Chromosomal location data frame
<code>Ref_start</code>	Starting position of deletion
<code>Ref_end</code>	Final position of deletion
<code>Chr</code>	Chromosome name

Value

Chromosomal location data frame with additional deletion info

Examples

```
gene_map = tugHall_dataset$gene_map
gene_map$pnts = ''
gm_new = add_deletion( gm = gene_map, Ref_start = 112775658, Ref_end = 112775716, Chr = '5')
# to check add_del ... add_dupl, please, use diff library and:
# diff_data( gm_ref, gm, show_unchanged_columns = TRUE, always_show_order = TRUE )
```

add_duplication	<i>Function to add duplication to gene map (chromosomal location data frame)</i>
-----------------	--

Description

Function to add duplication to gene map (chromosomal location data frame)

Usage

```
add_duplication(gm, Ref_start, Ref_end, Chr)
```

Arguments

gm	Chromosomal location data frame
Ref_start	Starting position of duplication
Ref_end	Final position of duplication
Chr	Chromosome name

Value

Chromosomal location data frame with additional duplication info

Examples

```
gene_map = tugHall_dataset$gene_map
gene_map$pnts = ''
gm_new = add_duplication( gm = gene_map, Ref_start = 112775658, Ref_end = 112775716, Chr = '5')
# to check add_del ... add_dupl, please, use diff library and:
# diff_data( gm_ref, gm, show_unchanged_columns = TRUE, always_show_order = TRUE )
```

add_pnt_mutation	<i>Function to add point mutation to data.frame gene_map (chromosomal location data frame)</i>
------------------	--

Description

Function to add point mutation to data.frame gene_map (chromosomal location data frame)

Usage

```
add_pnt_mutation(gm = gm, pos_pnt, Chr = Chr)
```

Arguments

gm	Chromosomal location data frame
pos_pnt	Position of point mutation
Chr	Chromosome name

Value

Chromosomal location data frame with additional point mutation info

Examples

```
gene_map = tugHall_dataset$gene_map
gene_map$pnts = ''
gm2 = add_pnt_mutation( gm = gene_map, pos_pnt = 112775637 , Chr = '5' )
```

calc_binom	<i>Function to calculate binomial distribution including BIG NUMBERS like 10^{12} and more using approximation with normal distribution</i>
------------	--

Description

Function to calculate binomial distribution including BIG NUMBERS like 10^{12} and more using approximation with normal distribution

Usage

```
calc_binom(tr, n, p)
```

Arguments

tr	Length of vector with successes trials
n	Number of independent Bernoulli trials
p	Probability to get successes in trials

Value

Vector of integer numbers of successes trials

Examples

```
calc_binom(tr = 3, n = 40, p = 0.9)
calc_binom(tr = 3, n = 4E20, p = 9E-9)
```

change_allele_A_by_cna	<i>Function to change copy number of the allele A of the point mutation at the allele B due to CNA</i>
------------------------	--

Description

Function to change copy number of the allele A of the point mutation at the allele B due to CNA

Usage

```
change_allele_A_by_cna(pnt1, start_end, t)
```

Arguments

pnt1 Object of class 'Point_Mutations'
 start_end Vector with initial and final positions of CNA
 t 'dup' or 'del' for duplication or deletion respectively

Value

NULL, but data of pnt1 is updated due to CNA

Examples

```
pnt1 = tugHall_dataset$pnt_clones[[ 2 ]] # pnt of allele A
start_end = c( pnt1$Phys_pos - 50 , pnt1$Phys_pos + 50 )
message( pnt1$Copy_number )
change_allele_A_by_cna( pnt1, start_end, t = 'dup' ) # View( pnt1 )
message( pnt1$Copy_number )
change_allele_A_by_cna( pnt1, start_end, t = 'del' ) # View( pnt1 )
message( pnt1$Copy_number )
```

change_pnt_by_cna	<i>Function to change the point mutation due to CNA</i>
-------------------	---

Description

Function to change the point mutation due to CNA

Usage

```
change_pnt_by_cna(pnt1, start_end, t)
```

Arguments

pnt1 Object of class 'Point_Mutations'
 start_end Vector with initial and final positions of CNA
 t 'dup' or 'del' for duplication or deletion respectively

Value

NULL, but pnt1 data is updated due to CNA

Examples

```
pnt1 = tugHall_dataset$pnt_clones[[ 1 ]]
start_end = c( pnt1$Phys_pos - 50 , pnt1$Phys_pos + 50 )
message( pnt1$Copy_number )
change_pnt_by_cna( pnt1, start_end, t = 'dup' ) # View( pnt1 )
message( pnt1$Copy_number )
change_pnt_by_cna( pnt1, start_end, t = 'del' ) # View( pnt1 )
message( pnt1$Copy_number )
```

check_pkg*Check the installation of a package for some functions*

Description

Check the installation of a package for some functions

Usage

```
check_pkg(pkg)
```

Arguments

pkg Package name

Value

if the package is installed then it returns NULL else it returns error message

Examples

```
check_pkg( pkg = 'grDevices' )
```

check_pnts*Function to check what pnts do fall into the range?*

Description

Function to check what pnts do fall into the range?

Usage

```
check_pnts(gm_w1)
```

Arguments

gm_w1 A row from data.frame gene_map

Value

Return the point mutations which fall into the range

Examples

```
gene_map = tugHall_dataset$gene_map
gene_map$pnts = ''
gene_map$pnts[6] = '112792451, 112792442'
gm_w1 = gene_map[6,]
check_pnts( gm_w1 )
```

chk_pnt_mut	<i>Function to check point mutations match or don't match into duplication or deletion</i>
-------------	--

Description

Function to check point mutations match or don't match into duplication or deletion

Usage

```
chk_pnt_mut(pnt1, Ref_start, Ref_end, Chr, prnt1)
```

Arguments

pnt1	Object of class 'Point_Mutations'
Ref_start	Initial position of CNA
Ref_end	Final position of CNA
Chr	Chromosome name
prnt1	Parental chromosome 1 or 2

Value

Logical: TRUE if point mutation matches CNA, FALSE if it doesn't match

Examples

```
pnt1 = tugHall_dataset$pnt_clones[[ 5 ]]
pstn = pnt1$Phys_pos[1]
message( pstn )
prnt1 = pnt1$Parental_1or2
Chr = pnt1$Chr
chk_pnt_mut( pnt1 , Ref_start = pstn - 200, Ref_end = pstn + 200, Chr, prnt1 )
chk_pnt_mut( pnt1 , Ref_start = pstn - 200, Ref_end = pstn - 100, Chr, prnt1 )
```

Clone-class	<i>Class 'Clone' for clones</i>
-------------	---------------------------------

Description

Class 'Clone' for clones

Fields

id	numeric. ID of a clone
parent	numeric. Parent ID (for first - 0)
N_cells	numeric. Number of cells in clone
c	numeric. Split counter as average value for all cells in clone
d	numeric. Probability of division

i numeric. Probability of Hayflick limit
 m numeric. Probability that gene normal function is destroyed due to epigenome abnormality / mutation rate
 a numeric. Probability of apoptosis for a cell in the clone
 s numeric. Coefficient in the sigmoid function of the mutation density
 k numeric. Probability of cell death by environment
 E numeric. Coefficient of friction term against to the split probability.
 Nmax numeric. Coefficient for determination the max number of cells that can exist in the primary tumor ($N_{max} = 1/E$)

 im numeric. Probability of the invasion/ metastatic transformation
 Ha numeric. Apoptosis hallmark value
 Him numeric. Invasion/ metastasis hallmark
 Hi numeric. Mitotic restriction hallmark (immortalization hallmark)
 Hd numeric. Growth/antigrowth hallmark (division rate hallmark)
 Hb numeric. Angiogenesis hallmark
 gene numeric. Vector of flags for each genes if they have driver mutation
 pasgene numeric. Vector of flags for each genes if they have passenger mutation
 PointMut_ID numeric. ID of point mutation in list of objects of class 'Point_Mutations'
 CNA_ID numeric. ID of CNA mutation in list of objects of class 'CNA_Mutations'
 mutden numeric. Gene mutation density
 invasion logical. Indicator that clone is metastatic (invasion/metastatic transformation occurred or not)
 primary logical. Logical variable is clone primary tumor or not (normal)
 birthday numeric. Time step of birth of clone

Examples

```

clone = tugHall_dataset$clones[[ 1 ]]
print(clone$Ha)
print(clone$N_cells)
clone$calcApoptosis() # to calculate apoptosis death probability based on mutation density

```

clone_copy

Function to make one copy for clone1 in clone_init function

Description

Function to make one copy for clone1 in clone_init function

Usage

```
clone_copy(clone1)
```

Arguments

clone1 Object of class 'Clone'

Value

New object of class 'Clone' with the same info and new ID

Examples

```
clone1 = tugHall_dataset$clones[[ 1 ]]
env = tugHall_dataset$env
define_parameters()
clone_copy(clone1)
```

CNA_Mutations-class *Class 'CNA_Mutations'*

Description

Class 'CNA_Mutations'

Fields

CNA_ID numeric. ID of CNA mutation

Parental_1or2 numeric. Parental chromosome, could be 1 or 2

dupOrdel character. dup for duplication or del for deletion

Chr character. Chromosome name

Ref_start numeric. Reference start position

Ref_end numeric. Reference final position

Gene_names character. Names of genes involved in CNA

MalfunctionedByCNA logical.

mut_order numeric. True for driver mutation and False for passenger mutation

Examples

```
cna = tugHall_dataset$cna_clones[[ 1 ]]
cna$save()    # to save as row of data.frame
cna$copy()
cna$initialize()
cna$show()    # After initialization
```

copy_CNA	<i>Function to copy CNA info</i>
----------	----------------------------------

Description

Function to copy CNA info

Usage

```
copy_CNA(CNA1)
```

Arguments

CNA1	Object of class 'CNA_Mutations'
------	---------------------------------

Value

The same object of class 'CNA_Mutations'

Examples

```
cna = tugHall_dataset$cna_clones[[ 1 ]]
cna2 = copy_CNA( cna )
cna$safe()
cna2$safe()
```

copy_files_to_Input	<i>Function to copy the files by default from extdata folder in the library to Input folder in the working directory</i>
---------------------	--

Description

Function to copy the files by default from extdata folder in the library to Input folder in the working directory

Usage

```
copy_files_to_Input(
  files = c("CCDS.current.txt", "CF.txt", "cloneinit.txt", "gene_hallmarks.txt",
    "gene_map.txt", "parameters.txt"),
  dir = "Input"
)
```

Arguments

files	Files to copy, vector of names of files by default: files = c('CCDS.current.txt', 'CF.txt', 'cloneinit.txt', 'gene_hallmarks.txt', 'gene_map.txt', 'parameter')
dir	Folder to where files should be save, by default dir = 'Input'

Value

List of logic numbers for each copied file, TRUE - success, FALSE - not success

Examples

```
files = c('CF.txt', 'cloneinit.txt', 'gene_hallmarks.txt', 'gene_map.txt', 'parameters.txt' )
copy_files_to_Input( files, dir = 'Input' )
```

copy_files_to_Output	<i>Function to copy the files of an example of simulation or from '/ext-data/Output/' folder in the library to '/Output/' folder in the working directory</i>
----------------------	---

Description

Function to copy the files of an example of simulation or from '/extdata/Output/' folder in the library to '/Output/' folder in the working directory

Usage

```
copy_files_to_Output(
  files = c("cloneout.txt", "CNA_mutations.txt", "point_mutations.txt", "gene_MAP.txt",
    "geneout.txt", "log.txt", "order_genes_dysfunction.txt", "VAF_data.txt", "VAF.txt",
    "weights.txt"),
  dir = "Output"
)
```

Arguments

files	Files to copy, vector of names of files by default: files = c('cloneout.txt', 'CNA_mutations.txt', 'point_mutations.txt', 'gene_MAP.txt', 'geneout.txt', 'log.txt', 'order_genes_dysfunction.txt', 'VAF_data.txt', 'VAF.txt', 'weights.txt')
dir	Folder to where files should be save, by default dir = 'Output'

Value

List of logic numbers for each copied file, TRUE - success, FALSE - not success

Examples

```
files = c( 'cloneout.txt', 'CNA_mutations.txt', 'point_mutations.txt', 'gene_MAP.txt' )
copy_files_to_Output( files )
files = c('geneout.txt', 'log.txt', 'VAF_data.txt', 'VAF.txt', 'weights.txt' )
copy_files_to_Output( files )
```

copy_pnt	<i>Function to copy of point mutation info</i>
----------	--

Description

Function to copy of point mutation info

Usage

```
copy_pnt(pnt1)
```

Arguments

pnt1 Object of class 'Point_Mutations'

Value

The same object of class 'Point_Mutations' with different ID

Examples

```
pnt = tugHall_dataset$pnt_clones[[ 1 ]]
copy_pnt( pnt ) # View( pnt )
```

copy_pnt_no_mutation	<i>Function to copy of pnt1 without mutation info for allele A</i>
----------------------	--

Description

Function to copy of pnt1 without mutation info for allele A

Usage

```
copy_pnt_no_mutation(pnt1)
```

Arguments

pnt1 Object of class 'Point_Mutations'

Value

Object of class 'Point_Mutations' for another chromosome

Examples

```
pnt = tugHall_dataset$pnt_clones[[ 1 ]]
copy_pnt_no_mutation( pnt ) # View( pnt )
```

```
define_compaction_factor
```

Define compaction factor

Description

Define compaction factor

Usage

```
define_compaction_factor(
  cf = data.frame(Ha = 1, Hb = 1, Hd = 1, Hi = 1, Him = 1),
  read_fl = TRUE,
  file_name = "./Input/CF.txt"
)
```

Arguments

cf	Data frame with compaction factors for all the hallmarks, for example, data.frame(Ha = 1, Hb = 1, Hd = 1, Hi = 1, Him = 1)
read_fl	Indicator to read file or not, logical type only
file_name	File name to read all the parameters, it is used only if read_fl == TRUE

Value

Data frame with with compaction factors for all the hallmarks

Examples

```
copy_files_to_Input()
define_compaction_factor( read_fl = TRUE , file_name = './Input/CF.txt' )
CF1 = CF
cf = data.frame( Ha = 0.1, Hb = 0.2, Hd = 0.7, Hi = 1, Him = 0.5 )
define_compaction_factor( cf = cf, read_fl = FALSE ) # View( c( CF, CF1 ) ) to compare
```

```
define_files_names
```

Function to define all the files names

Description

Function to define all the files names

Usage

```
define_files_names(
  mainDir = getwd(),
  sbdr_Input = "/Input",
  sbdr_Output = "/Output"
)
```

Arguments

mainDir	Working directory
sbdr_Input	Sub directory for input files
sbdr_Output	Sub directory for output files

Value

NULL, but all file names are defined in GLOBAL environment

Examples

```
define_files_names()
```

define_gene_location *Define genes' location in chromosome*

Description

Define genes' location in chromosome

Usage

```
define_gene_location(
  file_input = "Input/CCDS.current.txt",
  genes_list = c("CCDS4107.1", "CCDS8702.1", "CCDS43171.1", "CCDS11118.1")
)
```

Arguments

file_input	is a name of file to input where the information about genes location is defined. That is loaded from CCDS database https://ftp.ncbi.nlm.nih.gov/pub/CCDS/current_human/
genes_list	is a list of genes' names like CCDS4107.1 in the CCDS database.

Value

Function returns the table of genes' locations in DNA

Examples

```
copy_files_to_Input()
define_gene_location()
file_input = 'Input/CCDS.current.txt'
genes_list = c('CCDS4107.1', 'CCDS8702.1', 'CCDS43171.1', 'CCDS11118.1')
define_gene_location( file_input = file_input, genes_list = genes_list )
```

define_parameters	<i>Define all the parameters for a simulation</i>
-------------------	---

Description

Define all the parameters for a simulation

Usage

```
define_parameters(
  E0 = 1e-04,
  F0 = 10,
  m0 = 1e-07,
  uo = 0.9,
  us = 0.9,
  s0 = 10,
  k0 = 0.12,
  d0 = 0.4,
  censure_n = 10^5,
  censure_t = 50,
  m_dup = 1e-08,
  m_del = 1e-08,
  lambda_dup = 5000,
  lambda_del = 7000,
  uo_dup = 0.8,
  us_dup = 0.5,
  uo_del = 0,
  us_del = 0.8,
  CF = TRUE,
  model = c("proportional_metastatic", "threshold_metastatic", "simplified")[1],
  time_stop = 120,
  read_fl = FALSE,
  file_name = "../Input/parameters.txt",
  n_repeat = 1000,
  monitor = TRUE
)
```

Arguments

E0	Parameter in the division probability, numeric type only
F0	Parameter in the division probability, numeric type only
m0	Mutation probability for point mutation, numeric type only
uo	Oncogene mutation probability, numeric type only
us	Suppressor mutation probability, numeric type only
s0	Parameter in the sigmoid function, numeric type only
k0	Environmental death probability, numeric type only
d0	Initial probability to divide cells, numeric type only
censure_n	Max cell number where the program forcibly stops, integer type only

censore_t	Max time where the program forcibly stops, integer type only
m_dup	Mutation probability for duplication, numeric type only
m_del	Mutation probability for deletion, numeric type only
lambda_dup	CNA duplication average length (of the geometrical distribution for the length), integer type only
lambda_del	CNA deletion average length (of the geometrical distribution for the length), integer type only
uo_dup	Gene malfunction probability by CNA duplication for oncogene, numeric type only
us_dup	Gene malfunction probability by CNA duplication for suppressor, numeric type only
uo_del	Gene malfunction probability by CNA deletion for oncogene, numeric type only
us_del	Gene malfunction probability by CNA deletion for suppressor, numeric type only
CF	Compaction factor, logical type only. True means 'to use', False means 'do not use' Compaction factor for hallmarks variables
model	Name of the model to use. Can be 'proportional_metastatic' or 'threshold_metastatic' or 'simplified'
time_stop	Max time in seconds of running after that the program forcibly stops, integer type only
read_fl	Indicator to read file or not, logical type only
file_name	File name to read all the parameters, it is used only if read_fl == TRUE
n_repeat	Max number of repetition of the program until the NON-ZERO output will be getting, integer type only
monitor	The indicator to make monitor file during a simulation or do not make, logical type only

Value

Values of all the parameters

Examples

```
copy_files_to_Input()
define_parameters( read_fl = TRUE , file_name = './Input/parameters.txt' )
define_parameters( read_fl = FALSE )
```

define_par_for_plot *Function to change par for plots and after plotting it returns par values*

Description

Function to change par for plots and after plotting it returns par values

Usage

```
define_par_for_plot(change_par_back)
```

Arguments

`change_par_back`

logical. If TRUE it changes par options back after finishing a function call

Value

Change par() options and returns it (after finishing the function) to values before a function has started

Examples

```
define_par_for_plot( change_par_back = TRUE )
```

Environ-class

Class 'Environ'

Description

Class 'Environ'

Fields

T numeric. Time counter

N numeric. Number of normal cells

P numeric. Number of primary tumor cells

M numeric. Number of metastatic cells

F numeric. Coefficient that determines the maximal number of cells in pool of primary tumor cells

c numeric. Average number of divisions in pool of clones

d numeric. Mean value of splitting probability

i numeric. Average value of immortalization probability

a numeric. Average value of apoptosis probability

k numeric. Average probability of cell death via environment death

E numeric. Average value of coefficients of friction term

Nmax numeric. Maximal number of primary tumor cells that can exist in pool of clones

im numeric. Average value of invasion/metastasis probability

Ha numeric. Average value of apoptosis hallmark Ha

Him numeric. Average value of invasion/metastasis hallmark Him

Hi numeric. Average value of immortalization hallmark Hi

Hd numeric. Average value of growth/antigrowth hallmark Hd

Hb numeric. Average value of angiogenesis hallmark Hb

type numeric. Invasion / metastatic ratio

gene numeric. Cancer gene damage rate

mutden numeric. Average mutation rate

last_id numeric. Maximal ID in the pool of clones.

Examples

```
env = tugHall_dataset$env
print( env )
env$initFields()
```

generate_cna

*Function to generate object of CNA mutation***Description**

Function to generate object of CNA mutation

Usage

```
generate_cna(prnt1, genes, start_end, onco1, dupOrdel)
```

Arguments

prnt1	The 1st or 2nd parental chromosome
genes	Genes names
start_end	vector with start and final positions of CNA
onco1	Object of class 'OncoGene'
dupOrdel	It could be 'dup' or 'del' to denote duplication or deletion

Value

Object of class 'CNA_Mutations'

Examples

```
copy_files_to_Input()
define_parameters()
onco = tugHall_dataset$onco
gene_map = tugHall_dataset$gene_map
pnt_clones = tugHall_dataset$pnt_clones
cna_clones = tugHall_dataset$cna_clones
mut_order = 234
start_end = c(112775658, 112775716 )
generate_cna( prnt1 = 1, genes = 'APC', start_end = start_end, onco1, dupOrdel = 'dup' )
```

generate_pnt	<i>Function to generate an object of class 'Point_Mutations'</i>
--------------	--

Description

Function to generate an object of class 'Point_Mutations'

Usage

```
generate_pnt(prntl, gene, pos, onco1, Chr, mutation = NA)
```

Arguments

prntl	Parental chromosome, could be 1 or 2
gene	Gene name
pos	Position of point mutation
onco1	Object of class 'OncoGene'
Chr	Chromosome name
mutation	If mutation is NOT NA then MalfunctionedByPointMut = TRUE, else it is defined by corresponding probabilities

Value

Object of class 'Point_Mutations'

Examples

```
gm = tugHall_dataset$gene_map
gm_1_2 = list( gm, gm )
onco = tugHall_dataset$onco
pnt_clones = tugHall_dataset$pnt_clones
copy_files_to_Input()
define_parameters()
mut_order = 234 # As an example
pnt1 = generate_pnt( prntl = 1, gene = 'APC', pos = 112767192, onco, Chr = '5', mutation = NA )
```

generate_to_copy_pnt	<i>Function to generate the same object of class 'Point_Mutations' with coping all information from input object</i>
----------------------	--

Description

Function to generate the same object of class 'Point_Mutations' with coping all information from input object

Usage

```
generate_to_copy_pnt(pnt)
```

Arguments

pnt Object of class 'Point_Mutations'

Value

The same object of class 'Point_Mutations' with different ID

Examples

```
pnt = tugHall_dataset$pnt_clones[[ 1 ]]
pnt_clones = tugHall_dataset$pnt_clones
pnt2 = generate_to_copy_pnt( pnt )
```

gen_colors	<i>Function to make a large number of colors</i>
------------	--

Description

Function to make a large number of colors

Usage

```
gen_colors(nm = 12)
```

Arguments

nm Number of colors

Value

Vector of colors with length more than nm

Examples

```
clrs = gen_colors( nm = 120 )
```

get_cds_rna	<i>Function to get length of CDS and of genes from data.frame gene_map and related probabilities</i>
-------------	--

Description

Function to get length of CDS and of genes from data.frame gene_map and related probabilities

Usage

```
get_cds_rna(gm)
```

Arguments

gm data.frame gene_map with info about genes' location

Value

list(names, CDS, RNA, PROB, SUM, P0)

Examples

```
gene_map = tugHall_dataset$gene_map
define_parameters()
get_cds_rna( gm = gene_map )
```

get_cna_mutation	<i>Generation CNA mutation info</i>
------------------	-------------------------------------

Description

Generation CNA mutation info

Usage

```
get_cna_mutation(onco1, dupOrdel, gm_1_2)
```

Arguments

onco1	Object of class 'OncoGene'
dupOrdel	It could be 'dup' or 'del' to denote duplication or deletion
gm_1_2	List of two data frames (for 1st and 2nd parental chromosomes) with genes' location information

Value

List of (prntl - 1 or 2 parental chromosome, Chr - name of chromosome, genes - genes names, start_end - vector with start and end positions of CNA, w_cna - rows of CNA in gene_map data frame)

Examples

```
copy_files_to_Input()
define_parameters()
onco = tugHall_dataset$onco
gm = tugHall_dataset$gene_map
get_cna_mutation( onco1 = onco, dupOrdel = 'dup', gm_1_2 = list(gm, gm) )
get_cna_mutation( onco1 = onco, dupOrdel = 'del', gm_1_2 = list(gm, gm) )
```

get_flow_data	<i>Function to get data about last simulation from cloneoutfile</i>
---------------	---

Description

Function to get data about last simulation from cloneoutfile

Usage

```
get_flow_data(
  cloneoutfile,
  genefile,
  mainDir = getwd(),
  sbdr_Output = "/Output"
)
```

Arguments

cloneoutfile	Name of file to read data about clone evolution
genefile	Name of file with hallmarks values
mainDir	Working directory, by default mainDir = getwd()
sbdr_Output	Directory for output data getting from mainDir

Value

list of data.frames like onco, hall, data_last (data of last time step), data_avg (average data for all time steps), data_flow (data without average rows), time_max (max time step), pnt_mut and pnt_mut_B (data.frame of point mutations for both alleles and for allele B only) and cna_mut (data.frame of CNA mutations)

Examples

```
copy_files_to_Input()
define_files_names()
define_parameters( read_fl = TRUE , file_name = './Input/parameters.txt' )
define_compaction_factor( read_fl = TRUE , file_name = './Input/CF.txt' )
define_gene_location()
copy_files_to_Output()
dataset = get_flow_data(cloneoutfile, genefile, mainDir = getwd(), sbdr_Output = '/Output' )
# View(dataset)
```

get_len_cds_rna	<i>Function to get length of CDS and whole gene from gene_map data.frame</i>
-----------------	--

Description

Function to get length of CDS and whole gene from gene_map data.frame

Usage

```
get_len_cds_rna(gene_map)
```

Arguments

gene_map	data.frame with info about genes' locations
----------	---

Value

list of (Name, CDS, LEN_Genes) where Name is a vector of genes' names, CDS is a vector of CDS lengths, LEN_Genes is a vector of length of whole genes including introns and exons

Examples

```
gene_map = tugHall_dataset$gene_map
onco = tugHall_dataset$onco
get_len_cds_rna( gene_map)
```

get_order_of_genes_dysfunction	<i>Function to get order of genes' dysfunction</i>
--------------------------------	--

Description

Function to get order of genes' dysfunction

Usage

```
get_order_of_genes_dysfunction(
  pnt_mut,
  data_last,
  cna_mut,
  file_name = "../Output/order_genes_dysfunction.txt"
)
```

Arguments

pnt_mut	data.frame with info about all the point mutations
data_last	data.frame with data of simulation at the last time step
cna_mut	data.frame with info about all the CNA mutations
file_name	Name of file to save data

Value

data.frame of genes' dysfunction and save it in a file

Examples

```
pnt_mut = tugHall_dataset$pnt_mut
data_last = tugHall_dataset$data_last
cna_mut = tugHall_dataset$cna_mut
file_name = './Output/order_genes_dysfunction.txt'
rdr = get_order_of_genes_dysfunction( pnt_mut, data_last, cna_mut, file_name = file_name )
```

get_point_mutation	<i>Generation point mutation info</i>
--------------------	---------------------------------------

Description

Generation point mutation info

Usage

```
get_point_mutation(onco1, gm_1_2)
```

Arguments

onco1	Object of class 'OncoGene'
gm_1_2	List of two data frames (for 1st and 2nd parental chromosomes) with genes' location information

Value

list of (prntl - 1 or 2 parental chromosome, gene - gene name, pos - position of point mutation, Chr - name of chromosome)

Examples

```
gm = tugHall_dataset$gene_map
gm_1_2 = list( gm, gm )
onco = tugHall_dataset$onco
get_point_mutation( onco, gm_1_2 )
```

```
get_point_mutation_for_gene
```

Generation point mutation info for the particular gene

Description

Generation point mutation info for the particular gene

Usage

```
get_point_mutation_for_gene(onco1, gm_1_2, gene)
```

Arguments

onco1	Object of class 'OncoGene'
gm_1_2	List of two data frames (for 1st and 2nd parental chromosomes) with genes' location information
gene	Gene's name where point mutation should be occurred

Value

list of (prntl - 1 or 2 parental chromosome, gene - gene name, pos - position of point mutation, Chr - name of chromosome)

Examples

```
gm = tugHall_dataset$gene_map
gm_1_2 = list( gm, gm )
onco = tugHall_dataset$onco
get_point_mutation_for_gene( onco, gm_1_2, gene = 'APC')
get_point_mutation_for_gene( onco, gm_1_2, gene = 'KRAS')
```

```
get_rho_VAF
```

Function to get Variant allele frequencies (VAF) based on rho input parameters

Description

Function to get Variant allele frequencies (VAF) based on rho input parameters

Usage

```
get_rho_VAF(vf = NULL, rho = c(0, 0.1, 0.5), file_name = "./Output/VAF.txt")
```

Arguments

vf	data.frame getting from get_VAF() function
rho	Vector of rho parameter in the range (0,1)
file_name	Name of file to save VAF

Value

VAF for different rho with separation for metastatic cells and (primary tumor + speckled normal) cells

Examples

```
pnt_mut = tugHall_dataset$pnt_mut
data_last = tugHall_dataset$data_last
if ( !dir.exists('./Output') ) dir.create('./Output')
vf = get_VAF( pnt_mut, data_last, file_name = 'Output/VAF_data.txt')
VAF = get_rho_VAF( vf = vf, rho = c( 0.0, 0.1, 0.5 ) , file_name = './Output/VAF.txt' )
```

get_type

*Function to get type of the clone: normal, primary or metastatic***Description**

Function to get type of the clone: normal, primary or metastatic

Usage

```
get_type(clone1)
```

Arguments

clone1 Object of class 'Clone'

Value

One of characters 'normal', 'primary' or 'metastatic'

Examples

```
clone1 = tugHall_dataset$clones[[1]]
get_type( clone1 )
clone1 = tugHall_dataset$clones[[56]]
get_type( clone1 )
```

get_u_cna

*Function to choose probability of CNA mutation for several genes***Description**

Function to choose probability of CNA mutation for several genes

Usage

```
get_u_cna(genes, dupOrdel)
```

Arguments

genes	Names of genes, vector of names
dupOrdel	It could be 'dup' or 'del' to denote duplication or deletion

Value

Single value of maximal probability from probabilities for several genes

Examples

```
copy_files_to_Input()
define_parameters()
onco = tugHall_dataset$onco
get_u_cna( genes = 'APC', dupOrdel = 'dup' )
get_u_cna( genes = c('KRAS','APC'), dupOrdel = c('dup', 'del') )
```

get_VAF

Function to get data about Variant allele frequencies (VAF)

Description

Function to get data about Variant allele frequencies (VAF)

Usage

```
get_VAF(pnt_mut, data_last, file_name = "Output/VAF_data.txt")
```

Arguments

pnt_mut	data.frame with point mutation info
data_last	data.frame with data of simulation at the last time step
file_name	Name of file to save data

Value

data.frame with info about Variant allele frequencies

Examples

```
pnt_mut = tugHall_dataset$pnt_mut
data_last = tugHall_dataset$data_last
vf = get_VAF( pnt_mut, data_last, file_name = 'Output/VAF_data.txt')
```

HallMark-class	<i>Class 'HallMark'</i>
----------------	-------------------------

Description

Class 'HallMark'

Fields

Ha numeric. Apoptosis hallmark indexes of genes in onco\$name,
where onco is object of class OncoGene

Hi numeric. Immortalization hallmark indexes of genes in onco\$name,
where onco is object of class OncoGene

Hd numeric. Growth/antigrowth hallmark indexes of genes in onco\$name,
where onco is object of class OncoGene

Hb numeric. Angiogenesis hallmark indexes of genes in onco\$name,
where onco is object of class OncoGene

Him numeric. Invasion/metastatic transformation hallmark indexes of genes in onco\$name,
where onco is object of class OncoGene

Ha_w numeric. Apoptosis hallmark weights of genes

Hi_w numeric. Immortalization hallmark weights of genes

Hd_w numeric. Growth/antigrowth hallmark weights of genes

Hb_w numeric. Angiogenesis hallmark weights of genes

Him_w numeric. Invasion/metastatic transformation hallmark weights of genes

notHa numeric. Indexes of genes which are not in apoptosis hallmark

Examples

```
hall = tugHall_dataset$hall
print( hall )
hall$copy()
hall$show()
```

init_clones	<i>Function to read file with initial clones</i>
-------------	--

Description

Function to read file with initial clones

Usage

```
init_clones(clonefile, clone1)
```

Arguments

clonefile	File to read
clone1	Object of class 'Clone'

Value

List of objects of class 'Clone'

Examples

```
copy_files_to_Input()
define_files_names()
env = tugHall_dataset$env
define_parameters()
onco = tugHall_dataset$onco
clone1 = tugHall_dataset$clones[[ 1 ]]
clones = init_clones(clonefile, clone1) # View( clones )
```

init_onco_clones	<i>Function to make list of objects of class 'OncoGene' and generate initial onco settings for all clones (onco_clones)</i>
------------------	---

Description

Function to make list of objects of class 'OncoGene' and generate initial onco settings for all clones (onco_clones)

Usage

```
init_onco_clones(onco1, clones)
```

Arguments

onco1	Object of class 'OncoGene'
clones	List of objects of class 'Clone'

Value

List of objects of class 'OncoGene'

Examples

```
copy_files_to_Input()
define_files_names()
env = tugHall_dataset$env
define_parameters()
onco = tugHall_dataset$onco
clone1 = tugHall_dataset$clones[[ 1 ]]
clones = init_clones(clonefile, clone1) # View( clones )
onco_clones = init_onco_clones( onco1 = onco, clones )
```

init_pnt_clones	<i>Function to generate point mutations for initial clones</i>
-----------------	--

Description

Function to generate point mutations for initial clones

Usage

```
init_pnt_clones(clones, onco_clones)
```

Arguments

clones	List of objects of class 'Clone'
onco_clones	List of objects of class 'OncoGene'

Examples

```
clones = tugHall_dataset$clones
define_parameters()
onco = tugHall_dataset$onco
onco_clones = tugHall_dataset$onco_clones
copy_files_to_Input()
copy_files_to_Output()
define_gene_location()
pnt_clones = tugHall_dataset$pnt_clones
cna_clones = tugHall_dataset$cna_clones
mut_order = 234
## Not run:
init_pnt_clones( clones, onco_clones ) # change pnt_clones for initialization

## End(Not run)
```

make_map	<i>Function to make a gene_map data.frame with information of genes' locations</i>
----------	--

Description

Function to make a gene_map data.frame with information of genes' locations

Usage

```
make_map(
  f_out = "Input/map.txt",
  ls = c("CCDS4107.1", "CCDS8702.1", "CCDS43171.1", "CCDS11118.1"),
  f_in = "Input/CCDS.current.txt"
)
```

Arguments

f_out	Name of file to save gene_map data.frame
ls	List of IDs of genes corresponding CCDS database https://ftp.ncbi.nlm.nih.gov/pub/CCDS/current_h
f_in	Name of file to input downloaded from CCDS database

Value

gene_map data.frame with information of genes' locations for genes of interest

Examples

```
url = 'https://ftp.ncbi.nlm.nih.gov/pub/CCDS/current_human/CCDS.current.txt'
download.file( url = url, destfile = 'CCDS.current.txt')
ls = c( 'CCDS4107.1', 'CCDS8702.1', 'CCDS43171.1', 'CCDS11118.1' )
gene_map = make_map(f_out = 'map.txt', ls = ls, f_in = 'CCDS.current.txt' )
```

mixed_mut_order	<i>Function to get order of mutation for all possible types</i>
-----------------	---

Description

Function to get order of mutation for all possible types

Usage

```
mixed_mut_order(clone1)
```

Arguments

clone1	Object of class 'Clone'
--------	-------------------------

Value

data.frame with fields order, type, ID

Examples

```
clone = tugHall_dataset$clones[[ 46 ]]
clone$PointMut_ID
clone$CNA_ID
pnt_clones = tugHall_dataset$pnt_clones
cna_clones = tugHall_dataset$cna_clones
mixed_mut_order( clone )
```

model

Main function 'model' to simulate clones' evolution

Description

Main function 'model' to simulate clones' evolution

Usage

```
model(
  genefile,
  clonefile,
  geneoutfile,
  cloneoutfile,
  logoutfile,
  E0,
  F0,
  m0,
  uo,
  us,
  s0,
  k0,
  censure_n,
  censure_t,
  d0
)
```

Arguments

genefile	Name of file with input data of hallmarks
clonefile	Name of file with initial clones data
geneoutfile	Name of file with hallmark data
cloneoutfile	Name of file with cloneout data
logoutfile	File name of log file
E0	Parameter in the division probability, numeric type only
F0	Parameter in the division probability, numeric type only
m0	Mutation probability for point mutation, numeric type only
uo	Oncogene mutation probability, numeric type only
us	Suppressor mutation probability, numeric type only
s0	Parameter in the sigmoid function, numeric type only
k0	Environmental death probability, numeric type only
censure_n	Max cell number where the program forcibly stops, integer type only
censure_t	Max time where the program forcibly stops, integer type only
d0	Initial probability to divide cells, numeric type only

Value

List of (clones, onco_clones), where clones - list of objects of class 'Clone', and onco_clones - list of objects of class 'OncoGene'. During a simulation it saves data to geneoutfile.

Examples

```
copy_files_to_Input()
define_files_names()
define_gene_location()
define_parameters( read_fl = TRUE , file_name = './Input/parameters.txt' )
define_compaction_factor( read_fl = TRUE , file_name = './Input/CF.txt' )
time_stop = 3 # Duration of simulation time is 3 sec
## Not run:
res = model( genefile, clonefile, geneoutfile, cloneoutfile, logoutfile,
E0, F0, m0, uo, us, s0, k0, censore_n, censore_t, d0 )

## End(Not run)
```

modify_gene_map

Function to add the mutations to the data.frame gene_map

Description

Function to add the mutations to the data.frame gene_map

Usage

```
modify_gene_map(clone1, onco1)
```

Arguments

clone1	Object of class 'Clone'
onco1	Object of class 'OncoGene'

Value

list(gm1, gm2), where gm1 and gm2 are data.frames gene_maps with mutation information

Examples

```
clone = tugHall_dataset$clones[[ 46 ]]
onco = tugHall_dataset$onco_clones[[ 46 ]]
gene_map = tugHall_dataset$gene_map
pnt_clones = tugHall_dataset$pnt_clones
cna_clones = tugHall_dataset$cna_clones
gene_map$pnts = ''
## Not run:
gm_1_2 = modify_gene_map( clone , onco ) # View(gm_1_2)

## End(Not run)
```

number_N_P_M	<i>Function to get number of cells of a clone with indicator (normal, primary tumor or metastatic)</i>
--------------	--

Description

Function to get number of cells of a clone with indicator (normal, primary tumor or metastatic)

Usage

```
number_N_P_M(clone1)
```

Arguments

clone1 Object of class 'Clone'

Value

Vector c(N_normal, N_primary, N_metastatic)

Examples

```
clone1 = tugHall_dataset$clones[[ 1 ]]
number_N_P_M(clone1)
message( paste('Format is as follow: ', 'N_normal', 'N_primary', 'N_metastatic' ) )
```

OncoGene-class	<i>Class 'OncoGene'</i>
----------------	-------------------------

Description

Class 'OncoGene'

Fields

id numeric. ID is same as in clone (key for clones)

name character. Onco genes' names list

onsp character. Oncogene/suppressor indicator for each gene in list of names

len numeric. Lengths of onco genes

cds_1 numeric. Onco genes' CDS base lengths for parental chr 1

cds_2 numeric. Onco genes' CDS base lengths for parental chr 2

rna_1 numeric. Onco genes RNA base number length for parental chr 1 (exons+introns)

rna_2 numeric. Onco genes RNA base number length for parental chr 2 (exons+introns)

p0_1 numeric. Probability of absent of mutations for parental chr 1

p0_2 numeric. Probability of absent of mutations for parental chr 2

prob_1 numeric. Vector of relative probabilities for point mutation, deletion and duplication:

$$\text{prob} = \text{c}(m0 \times \text{sumCDS}, m_{\text{del}} \times \text{sumRNA}, m_{\text{dup}} \times \text{sumRNA}) / \text{sum}(m0 \times \text{sumCDS}, m_{\text{del}} \times \text{sumRNA}, m_{\text{dup}} \times \text{sumRNA})$$

```

prob_2 numeric.
sum_prob_1 numeric.
sum_prob_2 numeric.

```

Examples

```

onco = tugHall_dataset$onco
onco$copy()

```

onco_copy	<i>Function to make one copy for onco1 in init_onco_clones function</i>
-----------	---

Description

Function to make one copy for onco1 in init_onco_clones function

Usage

```
onco_copy(onco1)
```

Arguments

onco1 Object of class 'OncoGene'

Value

New object of class 'OncoGene' with the same info

Examples

```

onco1 = tugHall_dataset$onco_clones[[ 1 ]]
onco2 = onco_copy( onco1 ) # ID + 1

```

onco_update	<i>Function to update onco1 after mutation (for usage in trial_mutagenesis() function)</i>
-------------	--

Description

Function to update onco1 after mutation (for usage in trial_mutagenesis() function)

Usage

```
onco_update(onco1, gm)
```

Arguments

onco1 Object of class 'OncoGene'

gm data.frame gene_map

Value

onco1 with updated info

Examples

```
onco1 = tugHall_dataset$onco_clones[[ 1 ]]
copy_files_to_Input()
define_gene_location()
define_parameters()
onco_update( onco1, gm = list(gene_map, gene_map[1:42, ] ) ) # Check CDS length for TP53 gene
```

order_gene_map	<i>Function to order info in gene_map data.frame with information of genes' locations</i>
----------------	---

Description

Function to order info in gene_map data.frame with information of genes' locations

Usage

```
order_gene_map(gene_map)
```

Arguments

gene_map data.frame with information of genes' locations

Value

The same data.frame gene_map with ordered positions for each gene and each chromosome

Examples

```
gene_map = tugHall_dataset$gene_map
gene_map = order_gene_map( gene_map )
```

plot_2D	<i>Function to plot 2D figure of points $y = y(x)$</i>
---------	---

Description

Function to plot 2D figure of points $y = y(x)$

Usage

```
plot_2D(
  x,
  y,
  names = c("X", "Y"),
  pch = 18,
  col = "blue",
  cex = 1.2,
  xr = c(-10, 10),
  yr = c(-10, 10),
  safe_pdf = FALSE,
  filename = "./plot.pdf",
  change_par = TRUE
)
```

Arguments

x	Input data for axes X
y	Input data for axes Y
names	Vector of two characters with names for X and Y axes
pch	Parameter pch for plot function
col	Colors of points
cex	Parameter cex for plot function
xr	Range for X
yr	Range for Y
safe_pdf	Indicator to save plot to a file or not
filename	Name of file to save plot if safe_pdf == TRUE
change_par	Indicator to change par() or not for a plot. By default change_par = TRUE, after plot it will be returned to initial values

Value

NULL, making 2D plot using points

Examples

```
plot_2D( x=-5:5, y=-3:7 )
```

plot_2D_lines	<i>Function to plot 2D figure of lines $y_i = DF[, nl[i]]$, i - index</i>
---------------	--

Description

Function to plot 2D figure of lines $y_i = DF[, nl[i]]$, i - index

Usage

```

plot_2D_lines(
  x,
  DF,
  nl = 1:2,
  names = c("X", "Y"),
  legend_names = "",
  col = c("blue3", "darkmagenta", "red", "green4", "darkorange", "steelblue1"),
  cex = 1.2,
  lwd = 2,
  lt = c(1:6),
  xr = c(-10, 10),
  yr = c(-10, 10),
  safe_pdf = FALSE,
  filename = "./plot.pdf",
  type = "l",
  logscale = "",
  draw_key = TRUE,
  change_par = TRUE
)

```

Arguments

x	Input data for axes X
DF	data.frame with data to plot
nl	indexes of columns in DF to plot
names	Vector of two characters with names for X and Y axes
legend_names	Name of legend
col	Vector of colors for lines
cex	Parameter cex for plot function
lwd	Vector of width of lines
lt	Vector of types of lines
xr	Range for X
yr	Range for Y
safe_pdf	Indicator to save plot to a file or not
filename	Name of file to save plot if safe_pdf == TRUE
type	Parameter type in plot function
logscale	Parameter logscale in plot function
draw_key	Indicator to draw key or not
change_par	Indicator to change par() or not for a plot. By default change_par = TRUE, after plot it will be returned to initial values

Value

NULL, making 2D plot using lines

Examples

```

DF = tugHall_dataset$data_avg
plot_2D_lines( x = DF[, 1 ], DF, nl = 8:12 , xr = c(1,max(DF$Time) ), yr = c(0,1) )
xr = c(1,max(DF$Time) )
yr = c(0,max(DF[,14],DF[,16],DF[,17] ))
plot_2D_lines( x = DF[, 1 ], DF, nl = c(14,16,17) , xr =xr, yr = yr )
plot_2D_lines( x = DF[, 1 ], DF, nl = 18:22 , xr = c(1,max(DF$Time) ), yr = c(0,1) )

```

plot_average_simulation_data

Function to plot main data from data.frame with average data

Description

Function to plot main data from data.frame with average data

Usage

```
plot_average_simulation_data(data_avg, time_max)
```

Arguments

data_avg	data.frame with average values from cloneout.txt file
time_max	Maximal time step in a simulation

Value

NULL, draw many plot with average data

Examples

```

data_avg = tugHall_dataset$data_avg
time_max = tugHall_dataset$time_max
plot_average_simulation_data( data_avg , time_max = time_max )

```

plot_clone_evolution *Function to plot clone evolution*

Description

Function to plot clone evolution

Usage

```
plot_clone_evolution(
  data_flow,
  threshold = c(0.05, 1),
  lwd = 2,
  hue = c(" ", "random", "red", "orange", "yellow", "green", "blue", "purple", "pink",
    "monochrome")[1],
  luminosity = c(" ", "random", "light", "bright", "dark")[5],
  yr = NA,
  add_initial = TRUE,
  log_scale = FALSE,
  change_par = TRUE
)
```

Arguments

data_flow	data.frame with results of simulation at each time step
threshold	Vector two numbers from 0 to 1 to show clones with relative final numbers of cells in the range of threshold
lwd	Line width in the plot function
hue	Parameter hue in the function randomColor from library randomcoloR
luminosity	Parameter luminosity in the function randomColor from library randomcoloR
yr	Range for Y axes
add_initial	Indicator to add or do not add initial clones to plot
log_scale	Indicator to use log_scale or not for Y axes
change_par	Indicator to change par() or not for a plot. By default change_par = TRUE, after plot it will be returned to initial values

Value

NULL, making plot with clones evolution

Examples

```
data_flow = tugHall_dataset$data_flow
plot_clone_evolution( data_flow, threshold = c(0.01, 1 ), add_initial = TRUE, log_scale = FALSE )
plot_clone_evolution( data_flow, threshold = c(0, 0.01 ), add_initial = FALSE, log_scale = TRUE )
```

plot_order_dysfunction

Function to plot order of genes dysfunction as a step function with number of cells related to each order

Description

Function to plot order of genes dysfunction as a step function with number of cells related to each order

Usage

```
plot_order_dysfunction(
  rdr_dysf,
  pos = c(0, 100),
  logscale = "y",
  cex = 1,
  change_par = TRUE
)
```

Arguments

rdr_dysf	Order of genes dysfunction as a data.frame
pos	Coordinates of list of order of genes dysfunction
logscale	Parameter logscale for plot function
cex	Parameter cex for plot function
change_par	Indicator to change par() or not for a plot. By default change_par = TRUE, after plot it will be returned to initial values

Value

NULL, making plot with step function of order of genes' dysfunction

Examples

```
rdr_dysf = tugHall_dataset$rdr_dysf
plot_order_dysfunction( rdr_dysf , logscale = '', pos = c(8, 5000), cex = 1.4)
plot_order_dysfunction( rdr_dysf , logscale = 'y', pos = c(10, 100), cex = 1.2)
```

pnts_add_dlt

Function to subtract delta from position of point mutations

Description

Function to subtract delta from position of point mutations

Usage

```
pnts_add_dlt(gm_w1, dlt)
```

Arguments

gm_w1	A row from data.frame gene_map
dlt	Delta to subtract from positions of point mutations

Value

Return the pnts - dlt for one row of data.frame gene_map

Examples

```

gene_map = tugHall_dataset$gene_map
gene_map$pnts = ''
gene_map$pnts[6] = '112792451'
gm_w1 = gene_map[6,]
pnts_add_dlt( gm_w1 , dlt = 1000 )
pnts_add_dlt( gm_w1 , dlt = -1001 )

```

Point_Mutations-class *Class 'Point_Mutations'*

Description

Class 'Point_Mutations'

Fields

PointMut_ID numeric. ID of point mutation

Allele character. A or B allele

Parental_1or2 numeric. Parental chromosome, could be 1 or 2

Chr character. Chromosome name

Ref_pos numeric. Reference position

Phys_pos vector. Physical positions

Delta vector. Delta of positions

Copy_number numeric. Copy number of allele

Gene_name character. Gene's name

MalfunctionedByPointMut logical. True for driver mutation and False for passenger mutation

mut_order numeric. Number in order of mutation to reproduce the gene_map data.frame

Examples

```

pnt = tugHall_dataset$pnt_clones[[ 1 ]]
print( pnt )
pnt$copy()
pnt$show()
pnt$initialize()
pnt$show()
pnt = tugHall_dataset$pnt_clones[[ 3 ]]
pnt$save()    # save as row of data.frame

```

print_parameters	<i>Function to print GLOBAL parameters</i>
------------------	--

Description

Function to print GLOBAL parameters

Usage

```
print_parameters()
```

Value

Message with values of all the GLOBAL parameters

Examples

```
copy_files_to_Input()
define_parameters( read_fl = FALSE )
define_compaction_factor()
print_parameters()
```

read_file	<i>Function to read file</i>
-----------	------------------------------

Description

Function to read file

Usage

```
read_file(file_name = "", stringsAsFactors = FALSE, header = TRUE)
```

Arguments

file_name	Name of file to read
stringsAsFactors	Parameter for read.table function, by default stringsAsFactors = FALSE
header	Logical type to read or do not read head of a file

Value

data.frame of data from a file

Examples

```
f1 = system.file('extdata/Input', 'gene_map.txt', package = 'tugHall1.3', mustWork = TRUE )
read_file(file_name = f1, stringsAsFactors = FALSE )
f1 = system.file('extdata/Input', 'CF.txt', package = 'tugHall1.3', mustWork = TRUE )
read_file(file_name = f1, stringsAsFactors = FALSE, header = FALSE )
```

safe_pnt_mut	<i>Function to save 1 point mutation in a data frame</i>
--------------	--

Description

Function to save 1 point mutation in a data frame

Usage

```
safe_pnt_mut(pnt)
```

Arguments

pnt Object of class 'Point_Mutations'

Value

data frame with 1 row of point mutation info

Examples

```
pnt = tugHall_dataset$pnt_clones[[ 1 ]]
df = safe_pnt_mut( pnt ) # View( pnt )
```

simulation_example	<i>Example of simulation for lazy start</i>
--------------------	---

Description

Example of simulation for lazy start

Usage

```
simulation_example(verbose = TRUE, to_plot = TRUE, seed = NA)
```

Arguments

verbose Logical type to show or do not show messages during execution
to_plot Logical type to plot or do not plot graphical results of a simulation
seed Numeric type to set seed for a simulation, if seed = NA (by default) then it will be skipped

Value

List of results of simulation with default values for all the parameters

Examples

```
# it takes a time for a simulation and then it will demonstrates results, \cr
# so, please, wait for a while
simulation_example( verbose = FALSE , to_plot = FALSE )
```

sum_cell	<i>Aggregate data of a clone for environment object</i>
----------	---

Description

Aggregate data of a clone for environment object

Usage

```
sum_cell(env, clones)
```

Arguments

env	Object of class 'Environ'
clones	List of all the objects of class 'Clone'

Value

NULL, but global variable env is updated

Examples

```
clones = tugHall_dataset$clones
env = tugHall_dataset$env
sum_cell(env, clones)
message( paste0('Number of primary tumor cells in the pool of clones is ', env$P ) )
message( paste0('Number of normal cells in the pool of clones is ', env$N ) )
message( paste0('Number of metastatic cells in the pool of clones is ', env$M ) )
```

sum_mutation	<i>Serve function for sum_cell() function</i>
--------------	---

Description

Serve function for sum_cell() function

Usage

```
sum_mutation(clone1)
```

Arguments

clone1	Object of class 'Clone'
--------	-------------------------

Value

vector of clone1 variables to aggregate in sum_cell() function

Examples

```
clone1 = tugHall_dataset$clones[[ 1 ]]
sum_mutation(clone1)
```

sum_N_P_M

Function to calculate N and M numbers - normal and metastatic cells

Description

Function to calculate N and M numbers - normal and metastatic cells

Usage

```
sum_N_P_M(env, clones)
```

Arguments

env	Object of class 'Environ'
clones	List of all the objects of class 'Clone'

Value

Number of all the cells in a simulation (normal + primary tumor + metastatic)

Examples

```
clones = tugHall_dataset$clones
env = tugHall_dataset$env
env$M = 0
env$P = 0
env$N = 0 # View( env )
sum_N_P_M(env, clones) # View( env )
message( paste(env$N, env$P, env$M ) )
```

trial_complex

Function trial for complex case of models

Description

Function trial for complex case of models

Usage

```
trial_complex(clone1, onco1)
```

Arguments

clone1	Object of class 'Clone'
onco1	Object of class 'OncoGene'

Value

Number of new clones originated by clone1

Examples

```
clone1 = tugHall_dataset$clones[[ 1 ]]
onco1 = tugHall_dataset$onco
trial_complex( clone1, onco1 )
unlist( lapply( X = 1:20, FUN = function( x ) trial_complex( clone1, onco1 ) ) ) )
```

trial_mutagenesis	<i>Function for mutagenesis trial</i>
-------------------	---------------------------------------

Description

Function for mutagenesis trial

Usage

```
trial_mutagenesis(clone1, num_mut, onco1)
```

Arguments

clone1	Object of class 'Clone'
num_mut	Number of mutations in this NEW clone1
onco1	Object of class 'OncoGene' corresponding to clone1 (with the same ID)

Value

Changed object clone1, add related mutations to the lists of point mutations and/or CNA mutations

Examples

```
copy_files_to_Input()
copy_files_to_Output()
define_parameters()
clone1 = tugHall_dataset$clones[[ 1 ]]
onco1 = tugHall_dataset$onco_clones[[ 1 ]]
onco = tugHall_dataset$onco
define_gene_location()
pnt_clones = tugHall_dataset$pnt_clones
cna_clones = tugHall_dataset$cna_clones
mut_order = 234 # Just an example number
message( c('CNA mutation IDs ', paste(clone1$CNA_ID, collapse = ' ')) )
message( c('Point mutation IDs ', paste(clone1$PointMut_ID, collapse = ' ')) )
## Not run:
trial_mutagenesis( clone1, num_mut = 1, onco1 ) # it adds info to clone1
message( c('CNA mutation IDs ', paste(clone1$CNA_ID, collapse = ' ')) )
message( c('Point mutation IDs ', paste(clone1$PointMut_ID, collapse = ' ')) )

trial_mutagenesis( clone1, num_mut = 10, onco1 ) # it adds info to clone1
message( c('CNA mutation IDs ', paste(clone1$CNA_ID, collapse = ' ')) )
message( c('Point mutation IDs ', paste(clone1$PointMut_ID, collapse = ' ')) )

## End(Not run)
```

trial_simple	<i>Function trial for simplified case of model</i>
--------------	--

Description

Function trial for simplified case of model

Usage

```
trial_simple(clone1, onco1)
```

Arguments

clone1	Object of class 'Clone'
onco1	Object of class 'OncoGene'

Value

Number of new clones originated by clone1

Examples

```
clone1 = tugHall_dataset$clones[[ 1 ]]
onco1 = tugHall_dataset$onco
trial_simple( clone1, onco1 )
unlist( lapply( X = 1:20, FUN = function( x ) trial_simple( clone1, onco1 ) ) ) )
```

tugHall_dataset	<i>tugHall dataset named 'tugHall_dataset'</i>
-----------------	--

Description

Dataset contains all the necessary data.frames and objects to check functions of tugHall. Description of each data.frame and object could be found in documentation to tugHall package.

Usage

```
tugHall_dataset
```

Format

A data frame with 12 data.frames and 2 objects:

data_flow simulation data for all time steps, data from file cloneout.txt
data_last simulation data for the last time step, data from file cloneout.txt
data_avg simulation data averaged for the each time step, data from file cloneout.txt
pnt_mut data.frame with point mutation information
pnt_mut_B data.frame with point mutation information of mutated allele B
cna_mut data.frame with CNA mutation information

gene_map data.frame with genes' locations information

hall Object of class 'HallMark'

onco Object of class 'OncoGene'

time_max Value of maximal time step in an example simulation

CF Values of compaction factor

vf data.frame of preliminary data for VAF calculations

VAF_rho data.frame with VAF values for different rho

rdr_dysf data.frame of order of genes dysfunction for each clone

update_Hallmarks	<i>Function to update Hallmark and variable after division or under initialization</i>
------------------	--

Description

Function to update Hallmark and variable after division or under initialization

Usage

```
update_Hallmarks(clone1)
```

Arguments

clone1 Object of class 'Clone'

Value

The same object of class 'Clone' with updated fields

Examples

```
clone = tugHall_dataset$clones[[ 1 ]]
define_parameters()
hall = tugHall_dataset$hall
env = tugHall_dataset$env
update_Hallmarks( clone )
```

write_cloneout	<i>Function to write data to cloneout file at a time step</i>
----------------	---

Description

Function to write data to cloneout file at a time step

Usage

```
write_cloneout(outfile, env, clones, isFirst, onco_clones)
```

Arguments

outfile	File name for output info
env	Object of class 'Environ'
clones	List of objects of class 'Clone'
isFirst	logical type = TRUE as default
onco_clones	List of objects of class 'OncoGene'

Value

NULL, but add rows to output file with clone evolution data

Examples

```
env = tugHall_dataset$env
onco = tugHall_dataset$onco
if ( !dir.exists('./Output') ) dir.create('./Output')
clones = tugHall_dataset$clones
onco_clones = tugHall_dataset$onco_clones
write_header(outfile='./Output/exmpl.txt', env, onco)
write_cloneout( outfile = './Output/exmpl.txt', env, clones, isFirst = TRUE, onco_clones )
```

write_geneout	<i>Function to write info about HallMark data</i>
---------------	---

Description

Function to write info about HallMark data

Usage

```
write_geneout(outfile, hall, Compaction_factor, CF)
```

Arguments

outfile	File name for output info
hall	Object of class "HallMark"
Compaction_factor	Compaction factor, logical type only. True means 'to use', False means 'do not use' Compaction factor for hallmarks variables
CF	Vector with values of compaction factor for each hallmark

Value

NULL, but data will save to a file

Examples

```
copy_files_to_Input()
define_files_names()
define_parameters()
if ( !dir.exists('./Output') ) dir.create('./Output')
hall = tugHall_dataset$hall
onco = tugHall_dataset$onco
define_compaction_factor()
write_geneout(outfile = geneoutfile, hall, Compaction_factor, CF)
```

write_header	<i>Function to write the header to a file</i>
--------------	---

Description

Function to write the header to a file

Usage

```
write_header(outfile, env, onco)
```

Arguments

outfile	File name for output info
env	Object of class 'Environ'
onco	Object of class "OncoGene"

Value

NULL, but the header will save to a file and delete old info

Examples

```
env = tugHall_dataset$env
onco = tugHall_dataset$onco
if ( !dir.exists('./Output') ) dir.create('./Output')
write_header(outfile='./Output/exmpl.txt', env, onco)
```

write_log	<i>Function to write log file</i>
-----------	-----------------------------------

Description

Function to write log file

Usage

```
write_log(
  genefile,
  clonefile,
  geneoutfile,
  cloneoutfile,
  logoutfile,
  E0,
  F0,
  m0,
  uo,
  us,
  s0,
  k0,
  m_dup,
  m_del,
  lambda_dup,
  lambda_del,
  uo_dup,
  us_dup,
  uo_del,
  us_del,
  censore_n,
  censore_t,
  d0,
  Compaction_factor,
  model_name,
  time_stop,
  n_repeat,
  monitor
)
```

Arguments

genefile	File name of initial OncoGene information
clonefile	File name of info about initial clones
geneoutfile	File name for output info about OncoGene information
cloneoutfile	File name for output info with clone evolution data
logoutfile	Name of log file with all the parameters
E0	Parameter in the division probability, numeric type only
F0	Parameter in the division probability, numeric type only

m0	Mutation probability for point mutation, numeric type only
uo	Oncogene mutation probability, numeric type only
us	Suppressor mutation probability, numeric type only
s0	Parameter in the sigmoid function, numeric type only
k0	Environmental death probability, numeric type only
m_dup	Mutation probability for duplication, numeric type only
m_del	Mutation probability for deletion, numeric type only
lambda_dup	CNA duplication average length (of the geometrical distribution for the length), integer type only
lambda_del	CNA deletion average length (of the geometrical distribution for the length), integer type only
uo_dup	Gene malfunction probability by CNA duplication for oncogene, numeric type only
us_dup	Gene malfunction probability by CNA duplication for suppressor, numeric type only
uo_del	Gene malfunction probability by CNA deletion for oncogene, numeric type only
us_del	Gene malfunction probability by CNA deletion for suppressor, numeric type only
censore_n	Max cell number where the program forcibly stops, integer type only
censore_t	Max time where the program forcibly stops, integer type only
d0	Initial probability to divide cells, numeric type only
Compaction_factor	Compaction factor, logical type only. True means 'to use', False means 'do not use' Compaction factor for hallmarks variables
model_name	Name of the model to use. Can be 'proportional_metastatic' or 'threshold_metastatic' or 'simplified'
time_stop	Max time in seconds of running after that the program forcibly stops, integer type only
n_repeat	Max number of repetition of the program until the NON-ZERO output will be getting, integer type only
monitor	The indicator to make monitor file during a simulation or do not make, logical type only

Value

NULL, write log file to Output folder

Examples

```

copy_files_to_Input()
define_files_names()
define_parameters()
if ( !dir.exists('./Output') ) dir.create('./Output')
## Not run:
write_log(genefile, clonefile, geneoutfile, cloneoutfile, logoutfile,
E0, F0, m0, uo, us, s0, k0, m_dup, m_del, lambda_dup, lambda_del,
uo_dup, us_dup, uo_del, us_del, censore_n, censore_t, d0,
Compaction_factor, model_name, time_stop, n_repeat, monitor )

## End(Not run)

```

write_monitor	<i>Function to write a simulation monitoring data into the file_monitor</i>
---------------	---

Description

Function to write a simulation monitoring data into the file_monitor

Usage

```
write_monitor(outfile = file_monitor, start = FALSE, env, clones)
```

Arguments

outfile	File name for output info
start	Indicator to start from beginning (TRUE) or not (FALSE)
env	Object of class 'Environ'
clones	List of objects of class 'Clone'

Value

NULL, but info about current state of simulation will write to a file

Examples

```
env = tugHall_dataset$env
if ( !dir.exists('./Output') ) dir.create('./Output')
clones = tugHall_dataset$clones
onco_clones = tugHall_dataset$onco_clones
cna_clones = tugHall_dataset$cna_clones
pnt_clones = tugHall_dataset$pnt_clones
write_monitor( outfile = './Sim_monitoring.txt', start = TRUE , env, clones )
write_monitor( outfile = './Sim_monitoring.txt', start = FALSE , env, clones )
```

write_pnt_clones	<i>Function to write the point mutation info for all clones for all time steps, used at the last time step or after simulation</i>
------------------	--

Description

Function to write the point mutation info for all clones for all time steps, used at the last time step or after simulation

Usage

```
write_pnt_clones(pnt_clones, file_out = "Output/point_mutations.txt")
```

Arguments

pnt_clones	List of objects of class 'Point_Mutations'
file_out	File name to write

Value

NULL, but info will write to a file

Examples

```
pnt_clones = tugHall_dataset$pnt_clones
if ( !dir.exists('./Output') ) dir.create('./Output')
write_pnt_clones(pnt_clones, file_out = 'Output/point_mutations.txt')
```

write_weights	<i>Function to write info about relationship between genes and hallmarks</i>
---------------	--

Description

Function to write info about relationship between genes and hallmarks

Usage

```
write_weights(outfile, hall)
```

Arguments

outfile	File name for output info
hall	Object of class 'HallMark'

Value

NULL, but info about relationship between genes and hallmarks will write to a file

Examples

```
if ( !dir.exists('./Output') ) dir.create('./Output')
hall = tugHall_dataset$hall
onco = tugHall_dataset$onco
write_weights(outfile = './Output/weights.txt', hall)
```


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