# Package 'ActiveDriverWGS'

November 30, 2021

| n-<br>cu |
|----------|
|          |
|          |
|          |
|          |
|          |
|          |
|          |
|          |
|          |
|          |
|          |
|          |
|          |
|          |
|          |
|          |

2 .fix\_all\_results

|       | .get_signf_results   | 4  |
|-------|--|----|
|       | .make_mut_signatures   | 5  |
|       | .split_coord_fragments_in_BED  |    |
|       | ActiveDriverWGS  | 6  |
|       | ADWGS_test   | 8  |
|       | cancer_genes   | 10 |
|       | cancer_gene_sites  | 11 |
|       | cll_mutations  | 12 |
|       | format_muts  | 13 |
|       | prepare_elements_from_BED12  | 14 |
|       | prepare_elements_from_BED4   | 14 |
| Index |  | 16 |
| .fix_ | _all_results fix_all_results verifies that the results table has the correct format and p-values | d  |

## **Description**

fix\_all\_results verifies that the results table has the correct format and p-values

#### Usage

```
.fix_all_results(all_results)
```

## **Arguments**

all\_results

a data frame containing the following columns

id A string identifying the element of interest

**pp\_element** The p-value of the element

**element\_muts\_obs** The number of patients with a mutation in the element

**element\_muts\_exp** The expected number of patients with a mutation in the element with respect to background

**element\_enriched** A boolean indicating whether the element is enriched in mutations

**pp\_site** The p-value of the element

site\_muts\_obs The number of patients with a mutation in the site

**site\_muts\_exp** The expected number of patients with a mutation in the site with respect to element

site\_enriched A boolean indicating whether the site is enriched in mutations
result\_number A numeric indicator denoting the order in which the results
were calculated

## Value

the same data frame

```
.get_3n_context_of_mutations
```

This function finds the tri-nucleotide context of mutations

## **Description**

This function finds the tri-nucleotide context of mutations

## Usage

```
.get_3n_context_of_mutations(mutations, this_genome)
```

## **Arguments**

mutations A data frame with the following columns: chr, pos1, pos2, ref, alt, patient

chr autosomal chromosomes as chr1 to chr22 and sex chromosomes as chrX and chrY

**pos1** the start position of the mutation in base 1 coordinates

**pos2** the end position of the mutation in base 1 coordinates

**ref** the reference allele as a string containing the bases A, T, C or G **alt** the alternate allele as a string containing the bases A, T, C or G

patient the patient identifier as a string

this\_genome The reference genome object of BSgenome, for example BSgenome. Hsapiens. UCSC.hg19::Hsapiens

#### Value

A data frame consisting of the same columns as the original mutations data frame and sorted by SNVs and Indels with an additional column tag which indicates the trinucleotide context of the mutation

.get\_obs\_exp

Calculates the number of expected mutations based

## Description

Calculates the number of expected mutations based

## Usage

```
.get_obs_exp(hyp, select_positions, dfr, colname)
```

.get\_signf\_results

## **Arguments**

hyp hypothesis to be tested

select\_positions

boolean column which indicates which positions are in the element of interest

dfr a dataframe containing the data to be tested

colname name of the column which indicates the count of mutations in the positions of

interest

#### Value

a list of observed mutations and expected mutations

.get\_signf\_results

Returns significant results

## Description

Returns significant results

#### Usage

```
.get_signf_results(all_res)
```

## Arguments

all\_res a data frame containing the following columns

id A string identifying the element of interest

pp\_element The p-value of the element

element\_muts\_obs The number of patients with a mutation in the element

**element\_muts\_exp** The expected number of patients with a mutation in the element with respect to background

**element\_enriched** A boolean indicating whether the element is enriched in mutations

**pp\_site** The p-value of the element

site\_muts\_obs The number of patients with a mutation in the site

**site\_muts\_exp** The expected number of patients with a mutation in the site with respect to element

site\_enriched A boolean indicating whether the site is enriched in mutations

**result\_number** A numeric indicator denoting the order in which the results were calculated

.make\_mut\_signatures 5

## Value

the same data frame with three addition columns

fdr\_element The FDR corrected p-value of the element

fdr\_site The FDR corrected p-value of the site

has\_site\_mutations A V indicates the presence of site mutations

## **Description**

Makes mutational signatures

## Usage

```
.make_mut_signatures()
```

#### Value

a dataframe with mutational signatures

```
.split_coord_fragments_in_BED

Splits a BED12 file into separate regions
```

## Description

Splits a BED12 file into separate regions

## Usage

```
.split_coord_fragments_in_BED(i, coords)
```

## **Arguments**

i The i-th row of the coords data frame which needs to be split into separate

elements

coords The coords data frame which is the imported BED12 file

6 ActiveDriverWGS

#### Value

A data frame containing the following columns for a given BED12 identifier

chr autosomal chromosomes as chr1 to chr22 and sex chromosomes as chrX and chrY

**start** the start position of the element in base 0 coordinates (BED format)

end the end position of the element in base 0 coordinates (BED format)

id the element identifier - if the element contains multiple segments such as exons, each segment should be a separate row with the segment coordinates and the element identifier as id. Elements can be coding or noncoding such as exons of protein coding genes or active enhancers.

ActiveDriverWGS

ActiveDriverWGS is a driver discovery tool for simple somatic mutations in cancer whole genomes

## Description

ActiveDriverWGS is a driver discovery tool for simple somatic mutations in cancer whole genomes

## Usage

```
ActiveDriverWGS(
  mutations,
  elements,
  sites = NULL,
  window_size = 50000,
  filter_hyper_MB = 30,
  recovery.dir = NULL,
  mc.cores = 1,
  ref_genome = "hg19"
)
```

#### **Arguments**

mutations

A data frame containing the following columns: chr, pos1, pos2, ref, alt, patient.

chr autosomal chromosomes as chr1 to chr22 and sex chromosomes as chrX and chrY

**pos1** the start position of the mutation in base 1 coordinates

**pos2** the end position of the mutation in base 1 coordinates

ref the reference allele as a string containing the bases A, T, C, G or -

alt the alternate allele as a string containing the bases A, T, C, G or -

patient the patient identifier as a string

elements

A data frame containing the following columns: chr, start, end, id

chr autosomal chromosomes as chr1 to chr22 and sex chromosomes as chrX and chrY ActiveDriverWGS 7

**start** the start position of the element in base 0 coordinates (BED format) **end** the end position of the element in base 0 coordinates (BED format)

id the element identifier - if the element contains multiple segments such as exons, each segment should be a separate row with the segment coordinates and the element identifier as id. Elements can be coding or noncoding such as exons of protein coding genes or active enhancers.

sites A data frame containing the following columns: chr, start, end, id

chr autosomal chromosomes as chr1 to chr22 and sex chromosomes as chrX and chrY

start the start position of the site in base 0 coordinates (BED format)

end the end position of the site in base 0 coordinates (BED format)

id the identifier of the element. id's need to match with those listed in the object elements.

window\_size An integer indicating the size of the background window in base pairs that is

used to establish the expected mutation rate and respective null model. The

default is 50000bps

filter\_hyper\_MB

Hyper-mutated samples carry many passenger mutations and dilute the signal of true drivers. Samples with a rate greater than filter\_hyper\_MB mutations per megabase are excluded. The default is 30 mutations per megabase.

recovery.dir The directory for storing recovery files. If the directory does not exist, Ac-

tiveDriverWGS will create the directory. If the parameter is unspecified, recovery files will not be saved. As an ActiveDriverWGS query for large datasets may be computationally heavy, specifying a recovery directory will recover pre-

viously computed results if a query is interrupted.

mc.cores The number of cores which can be used if multiple cores are available. The

default is 1.

ref\_genome The reference genome used on the analysis. The default option is "hg19", other

options are "hg38", "mm9" and "mm10".

#### Value

A data frame containing the results of driver discovery containing the following columns: id, pp\_element, element\_muts\_obs, element\_muts\_exp, element\_enriched, pp\_site, site\_muts\_obs, site\_muts\_exp, site\_enriched, fdr\_element, fdr\_site

id A string identifying the element of interest

**pp\_element** The p-value of the element

**element muts obs** The number of patients with a mutation in the element

element\_muts\_exp The expected number of patients with a mutation in the element with respect
to background

**element enriched** A boolean indicating whether the element is enriched in mutations

**pp\_site** The p-value of the site

**site\_muts\_obs** The number of patients with a mutation in the site

8 ADWGS\_test

site\_muts\_exp The expected number of patients with a mutation in the site with respect to elementsite\_enriched A boolean indicating whether the site is enriched in mutations

fdr\_element The FDR corrected p-value of the element

fdr\_site The FDR corrected p-value of the site

has\_site\_mutations A V indicates the presence of site mutations

## **Examples**

```
data(cancer_genes)
data(cll_mutations)

some_genes = c("ATM", "MYD88", "NOTCH1", "SF3B1", "XPO1",
   "SOCS1", "CNOT3", "DDX3X", "KMT2A", "HIF1A", "APC")

result = ActiveDriverWGS(mutations = cll_mutations,
elements = cancer_genes[cancer_genes$id %in% some_genes,])
```

ADWGS\_test

ADWGS\_test executes the statistical test for ActiveDriverWGS

## **Description**

ADWGS\_test executes the statistical test for ActiveDriverWGS

## Usage

```
ADWGS_test(
   id,
   gr_element_coords,
   gr_site_coords,
   gr_maf,
   win_size,
   this_genome
)
```

## **Arguments**

id

A string used to identify the element of interest. id corresponds to an element in the id column of the elements file

```
gr_element_coords
```

A GenomicRanges object that describes the elements of interest containing the chromosome, start and end coordinates, and an mcols column corresponding to id

ADWGS\_test 9

gr\_site\_coords A GenomicRanges object that describes the sites of interest which reside in the

elements of interest containing the chromosome, start and end coordinates, and an mcols column corresponding to id. Examples of sites include transcription factor binding sites in promoter regions or phosphosites in exons of protein coding genes. An empty GenomicRanges object nullifies the requirement for sites

to exist.

gr\_maf A GenomicRanges object that describes the mutations in the dataset containing

the chromosome, start and end coordinates, patient id, and trinucleotide context

win\_size An integer indicating the size of the background window in base pairs that is

used to establish the expected mutation rate and respective null model. The

default is 50000bps

this\_genome The reference genome object of BSgenome, for example BSgenome. Hsapiens. UCSC.hg19::Hsapiens

#### Value

A data frame containing the following columns

id A string identifying the element of interest

**pp\_element** The p-value of the element

element\_muts\_obs The number of patients with a mutation in the element

**element\_muts\_exp** The expected number of patients with a mutation in the element with respect to background

**element\_enriched** A boolean indicating whether the element is enriched in mutations

**pp\_site** The p-value of the site

**site\_muts\_obs** The number of patients with a mutation in the site

site\_muts\_exp The expected number of patients with a mutation in the site with respect to element

site\_enriched A boolean indicating whether the site is enriched in mutations

result\_number A numeric indicator denoting the order in which the results were calculated

fdr\_element The FDR corrected p-value of the element

fdr\_site The FDR corrected p-value of the site

has\_site\_mutations A V indicates the presence of site mutations

```
library(GenomicRanges)

# Regions
data(cancer_genes)
gr_element_coords = GRanges(seqnames = cancer_genes$chr,
IRanges(start = cancer_genes$start, end = cancer_genes$end),
mcols = cancer_genes$id)

# Sites (NULL)
gr_site_coords = GRanges(c(seqnames=NULL,ranges=NULL,strand=NULL))
```

10 cancer\_genes

```
# Reference genome
this_genome = BSgenome.Hsapiens.UCSC.hg19::Hsapiens

# Mutations
data(cll_mutations)
cll_mutations = format_muts(cll_mutations, this_genome = this_genome)
gr_maf = GRanges(cll_mutations$chr,
IRanges(cll_mutations$pos1, cll_mutations$pos2),
mcols=cll_mutations[,c("patient", "tag")])

# ADWGS_test
id = "ATM"
result = ADWGS_test(id, gr_element_coords, gr_site_coords, gr_maf,
win_size = 50000, this_genome = this_genome)
```

cancer\_genes

cancer\_genes

## **Description**

protein coding genes from gencode v.19, cancer genes adapted from the Cancer Gene Census (November, 2018). Genes affected solely by amplifications, deletions and translations were removed.

#### Usage

```
data(cancer_genes)
```

## Format

A data frame containing the following columns: chr, start, end, id

chr autosomal chromosomes as chr1 to chr22 and sex chromosomes as chrX and chrY

**start** the start position of the element in base 0 coordinates (BED format)

**end** the end position of the element in base 0 coordinates (BED format)

id the element identifier - if the element contains multiple segments such as exons, each segment should be a separate row with the segment coordinates and the element identifier as id. Elements can be coding or noncoding such as exons of protein coding genes or active enhancers.

#### Source

#### **GENCODE**

#### References

Harrow, Jennifer, et al. "GENCODE: the reference human genome annotation for The ENCODE Project." Genome research 22.9 (2012): 1760-1774. (PubMed)

cancer\_gene\_sites 11

## **Examples**

```
data(cancer_genes)

data(cll_mutations)
ActiveDriverWGS(mutations = cll_mutations, elements = cancer_genes)
```

cancer\_gene\_sites

post-translational modification sites found in cancer genes

## **Description**

post-translational modification sites found in cancer genes

## Usage

```
data(cancer_gene_sites)
```

#### **Format**

A data frame containing the following columns: chr, start, end, id

chr autosomal chromosomes as chr1 to chr22 and sex chromosomes as chrX and chrY

**start** the start position of the site in base 0 coordinates (BED format)

**end** the end position of the site in base 0 coordinates (BED format)

id the site identifier - each site should contain only 1 segment and a unique id. If ids are duplicated, each segment of the site will be treated as an individual site. Sites can be coding or noncoding such as phosphosites of protein coding genes in genomic coordinates or transcription factor binding sites of active enhancers.

#### Source

#### PubMed

## References

Wadi, Lina, et al. "ActiveDriverDB: human disease mutations and genome variation in post-translational modification sites of proteins." Nucleic Acids Res. (2018): Jan 4;46(D1):D901-D910. (PubMed)

```
data(cancer_gene_sites)

data(cll_mutations)
data(cancer_genes)
ActiveDriverWGS(mutations = cll_mutations, elements = cancer_genes, sites = cancer_gene_sites)
```

12 cll\_mutations

cll\_mutations

CLL mutations

## **Description**

CLL whole genome simple somatic mutations from Alexandrov et, 2013

## Usage

```
data(cll_mutations)
```

## **Format**

A data frame containing the following columns: chr, pos1, pos2, ref, alt, patient.

chr autosomal chromosomes as chr1 to chr22 and sex chromosomes as chrX and chrY

**pos1** the start position of the mutation in base 1 coordinates

pos2 the end position of the mutation in base 1 coordinates

ref the reference allele as a string containing the bases A, T, C or G

alt the alternate allele as a string containing the bases A, T, C or G

patient the patient identifier as a string

## Source

**Publication** 

#### References

Alexandrov, Ludmil B., et al. "Signatures of mutational processes in human cancer." Nature 500.7463 (2013): 415. (PubMed)

```
data(cll_mutations)

data(cancer_genes)
ActiveDriverWGS(mutations = cll_mutations, elements = cancer_genes)
```

format\_muts 13

| format_muts | This function filters hypermutated samples and returns the formatted mutations with the appropriate trinucleotide context |
|-------------|---|
|             |   |

## **Description**

This function filters hypermutated samples and returns the formatted mutations with the appropriate trinucleotide context

## Usage

```
format_muts(mutations, this_genome, filter_hyper_MB = NA)
```

## **Arguments**

mutations A data frame with the following columns: chr, pos1, pos2, ref, alt, patient

chr autosomal chromosomes as chr1 to chr22 and sex chromosomes as chrX and chrY

**pos1** the start position of the mutation in base 1 coordinates

pos2 the end position of the mutation in base 1 coordinates

ref the reference allele as a string containing the bases A, T, C or G

alt the alternate allele as a string containing the bases A, T, C or G

patient the patient identifier as a string

this\_genome The reference genome object of BSgenome

filter\_hyper\_MB

The number of mutations per megabase for which a sample is considered hypermutated. Hypermutated samples will be removed in further analyses.

## Value

a data frame called mutations which has been formatted with an extra column for trinucleotide context

```
data(cll_mutations)
this_genome = BSgenome.Hsapiens.UCSC.hg19::Hsapiens
formatted_mutations = format_muts(cll_mutations[1:10,],
filter_hyper_MB = 30, this_genome = this_genome)
```

```
prepare_elements_from_BED12
```

Prepares element coords from a BED12 file

## Description

Prepares element coords from a BED12 file

## Usage

```
prepare_elements_from_BED12(fname)
```

## Arguments

fname

The file name of a BED12 file containing the desired elements. For further documentation on the BED12 format, refer to the UCSC website.

#### Value

A data frame containing the following columns to be used as the input element coords to ActiveDriverWGS

chr autosomal chromosomes as chr1 to chr22 and sex chromosomes as chrX and chrY

**start** the start position of the element in base 0 coordinates (BED format)

end the end position of the element in base 0 coordinates (BED format)

id the element identifier - if the element contains multiple segments such as exons, each segment should be a separate row with the segment coordinates and the element identifier as id. Elements can be coding or noncoding such as exons of protein coding genes or active enhancers.

## **Examples**

```
elements = prepare_elements_from_BED12(system.file("extdata",
   "chr17.coding_regions.bed",
   package = "ActiveDriverWGS",
   mustWork = TRUE))
```

```
prepare_elements_from_BED4
```

Prepares element coords from a BED4 file

## Description

Prepares element coords from a BED4 file

## Usage

```
prepare_elements_from_BED4(fname)
```

## **Arguments**

fname

The file name of a BED4 file containing the desired elements. For further documentation on the BED4 format, refer to the UCSC website.

## Value

A data frame containing the following columns to be used as the input element coords to ActiveDriverWGS

chr autosomal chromosomes as chr1 to chr22 and sex chromosomes as chrX and chrY

start the start position of the element in base 0 coordinates (BED format)

**end** the end position of the element in base 0 coordinates (BED format)

id the element identifier - if the element contains multiple segments such as exons, each segment should be a separate row with the segment coordinates and the element identifier as id. Elements can be coding or noncoding such as exons of protein coding genes or active enhancers.

```
elements = prepare_elements_from_BED4(system.file("extdata",
"mini.ptm.bed",
package = "ActiveDriverWGS",
mustWork = TRUE))
```

## **Index**

```
* datasets
    cancer_gene_sites, 11
    cancer_genes, 10
    cll_mutations, 12
.fix_all_results, 2
. \verb"get_3n_context_of_mutations", 3
.get_obs_exp, 3
.get\_signf\_results, 4
.make\_mut\_signatures, 5
.split\_coord\_fragments\_in\_BED, 5
ActiveDriverWGS, 6
ADWGS_test, 8
cancer\_gene\_sites, 11
cancer_genes, 10
\verb|cll_mutations|, 12|
format_muts, 13
prepare_elements_from_BED12, 14
prepare_elements_from_BED4, 14
```