Package 'Uniquorn'

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Title Identification of cancer cell lines based on their weighted mutational/ variational fingerprint

Version 2.29.2

Description 'Uniquorn' enables users to identify cancer cell lines.

Cancer cell line misidentification and cross-

contamination reprents a significant challenge for cancer researchers.

The identification is vital and in the frame of this package based on the locations/ loci of somatic and germline mutations/ variations.

The input format is vcf/ vcf.gz and the files have to contain a single cancer cell line sample (i.e. a single member/genotype/gt column in the vcf file).

Imports stringr, R.utils, WriteXLS, stats, doParallel, foreach, GenomicRanges, IRanges, VariantAnnotation, data.table

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```

add_custom_vcf_to_database This function adds the variants of parsed custom CCLs to a monet DB instance

Description

add_custom_vcf_to_database This function adds the variants of parsed custom CCLs to a monet DB instance

```
add_custom_vcf_to_database(
    vcf_input_files,
    ref_gen = "GRCH37",
    library_name = "CUSTOM",
    n_threads = 1,
    test_mode = FALSE
)
```

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Arguments

vcf_input_files

a character vector containing the input vcf files. This may be one or many vcf

files.

ref_gen a character string specifying the reference genome version. All training sets are

associated with a reference genome version. Default is "GRCH37".

library_name a character string giving the name of the library to add the cancer cell lines to.

Default is "CUSTOM". Library name will be automatically added as a suffix to

the identifier.

n_threads an integer specifying the number of threads to be used.

test_mode Is this a test? Just for internal use

Value

Message wheather the adding was successful

Examples

```
HT29_vcf_file = system.file("extdata/HT29_TEST.vcf", package = "Uniquorn");
add_custom_vcf_to_database(
    vcf_input_files = HT29_vcf_file,
    library_name = "CELLMINER",
    ref_gen = "GRCH37",
    n_threads = 1,
    test_mode = TRUE
)
```

add_missing_cls

add_missing_cls

Description

```
add_missing_cls
```

Usage

```
add_missing_cls(res_table, dif_cls)
```

Arguments

res_table Table that contains the identification results

dif_cls Missing CLs

Value

Results table with added missing cls

```
add\_penalty\_statistics \\ add\_penalty\_statistics
```

Add penalty statistics to results

Usage

```
add_penalty_statistics(match_t, minimum_matching_mutations)
```

Arguments

```
match_t object that contains the matching variants
minimum_matching_mutations
a numerical giving the minimum amount of mutations that has to match between
query and training sample for a positive prediction
```

Value

The updated statistics

```
add\_p\_q\_values\_statistics \\ add\_p\_q\_values\_statistics
```

Description

A hypergeometric distribution-assumption allows to calculate the p-values for a significant or non-significant overlap in this function

```
add_p_q_values_statistics(
   g_query,
   match_t,
   p_value,
   ref_gen,
   minimum_matching_mutations,
   top_hits_per_library
)
```

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Arguments

	g_query	IRanges object that contains the query variants
	match_t	A table that contains the nubmber of matching variants
	p_value	Threshold for the significance p-value
	ref_gen	Reference genome version
minimum_matching_mutations		
		Manual lower amount of matching mutations require for a significant match
		between a query and a reference
top_hits_per_library		
		limits significant similarities to the first n hits

Details

```
add_p_q_values_statistics Calculates the p-values
```

Value

R table with a statistic

Description

Creates BED files from the found and not found annotated mutations

Usage

```
create_bed_file(
match_t,
vcf_fingerprint,
output_file,
ref_gen,
manual_identifier
)
```

Arguments

match_t R table which contains the mutations from the training database for the cancer cell lines

vcf_fingerprint contains the mutations that are present in the query cancer cell line's vcf file

output_file Path to output file

ref_gen Reference genome version

manual_identifier

Manually enter a vector of CL name(s) whose bed files should be created, independently from them passing the detection threshold

6 identify_vcf_file

Value

Returns a message which indicates if the BED file creation has succeeded

Description

Identifies a cancer cell lines contained in a vcf file based on the pattern (start & length) of all contained mutations/ variations.

Usage

```
identify_vcf_file(
    vcf_file,
   output_file,
   ref_gen,
   minimum_matching_mutations,
   mutational_weight_inclusion_threshold,
   write_xls,
    output_bed_file,
    top_hits_per_library,
    manual_identifier,
    verbose,
   p_value,
   confidence_score,
   n_threads,
   write_results
)
```

Arguments

write_xls

vcf_file	Input vcf file. Only one sample column allowed.	
output_file	Path of the output file. If blank, autogenerated as name of input file plus '_uniquorn_ident.tab' suffix.	
ref_gen	Reference genome version. All training sets are associated with a reference genome version. Default: GRCH37	
minimum_matching_mutations		
	The minimum amount of mutations that has to match between query and training sample for a positive prediction	
mutational_weight_inclusion_threshold		
	Include only mutations with a weight of at least x. Range: 0.0 to 1.0 . 1= unique to CL. ~ 0 = found in many CL samples.	

Create identification results additionally as xls file for easier reading

output_bed_file

If BED files for IGV visualization should be created for the Cancer Cell lines that pass the threshold

top_hits_per_library

Limit the number of significant similarities per library to n (default 3) many hits. Is particularly used in contexts when heterogeneous query and reference CCLs are being compared.

manual_identifier

Manually enter a vector of CL name(s) whose bed files should be created, independently from them passing the detection threshold

verbose Print additional information

p_value Required p-value for identification. Note that if you set the confidence score,

the confidence score overrides the p-value

confidence_score

Cutoff for positive prediction between 0 and 100. Calculated by transforming the p-value by -1 * log(p-value) Note that if you set the confidence score, the

confidence score overrides the p-value

n_threads Number of threads to be used write_results Write identification results to file

Details

identify_vcf_file parses the vcf file and predicts the identity of the sample

Value

R table with a statistic of the identification result

Examples

```
HT29_vcf_file = system.file("extdata/HT29.vcf", package = "Uniquorn");
identification = identify_vcf_file(
    vcf_file = HT29_vcf_file,
    verbose = FALSE,
    write_results = FALSE
)
```

initiate_canonical_databases

initiate_canonical_databases

Description

Parses data into r list variable

Usage

```
initiate_canonical_databases(
    cosmic_file = "CosmicCLP_MutantExport.tsv.gz",
    ccle_file = "CCLE_mutations.csv",
    ccle_sample_file = "sample_info.csv",
    ref_gen = "GRCH38"
)
```

Arguments

```
The path to the Cosmic CLP file. The Cosmic file can be obtained from "https://cancer.sanger.ac.uk/cell_li and should be labeled "CosmicCLP_MutantExport.tsv.gz". Ensure that the right reference genome is used

CCle_file The path to the ccle DNA genotype data file. It should be labeled "CCLE_mutations.csv".

Ensure that the right reference genome is used

Ccle_sample_file

The path to the CCLE sample file. It should be labeled "sample_info.csv" containing both the DepMap ID and corresponding cell line name.

Reference genome version
```

Value

Returns message if parsing process has succeeded

Examples

```
initiate_canonical_databases(
    cosmic_file = "CosmicCLP_MutantExport.tsv.gz",
    ccle_file = "CCLE_mutations.csv",
    ccle_sample_file = "sample_info.csv",
    ref_gen = "GRCH38"
)
```

```
init\_and\_load\_identification \\ init\_and\_load\_identification
```

Description

Initiate the analysis Output basic information

Usage

```
init_and_load_identification(
    verbose,
    ref_gen,
    vcf_file,
    output_dir
)
```

Arguments

verbose Print additional information

ref_gen Reference genome version. All training sets are associated with a reference genome version. Default: GRCH37

vcf_file Path to vcf_file

output_dir Output directory for identification results

Details

init_and_load_identification parses vcf file and output basic information

Value

Three file path instances and the fingerprint

Description

Matches query ccl to the database

```
match_query_ccl_to_database(
   g_query,
   ref_gen = "GRCH37",
   library_name,
   mutational_weight_inclusion_threshold
)
```

Arguments

g_query IRanges object that contains the variants

ref_gen Reference genome version. All training sets are associated with a reference

genome version. Default: GRCH37

library_name a character string giving the name of the library

mutational_weight_inclusion_threshold

a numerical giving the lower bound for mutational weight to be included

Value

The R Table sim_list which contains the CoSMIC CLP fingerprints

Description

Parses ccle genotype data

Usage

```
parse_ccle_genotype_data(ccle_file, ccle_sample_file, ref_gen = "GRCH38")
```

Arguments

ccle_file Path to CCLE file on hard disk

ccle_sample_file

Path to CCLE sample file

ref_gen Reference genome version

Value

The R Table sim_list which contains the CCLE fingerprints

Parses cosmic genotype data

Usage

```
parse_cosmic_genotype_data(cosmic_file, ref_gen = "GRCH38")
```

Arguments

```
cosmic_file Path to cosmic clp file in hard disk ref_gen Reference genome version
```

Value

The R Table sim_list which contains the CoSMIC CLP fingerprints

```
parse_vcf_file Filter Parsed VCF Files
```

Description

Intern utility function. Filters the parsed VCF file for all informations except for the start and length of variations/mutations.

Usage

```
parse_vcf_file(
    vcf_file,
    ref_gen,
    library_name
)
```

Arguments

vcf_file character string giving the path to the vcf file on the operating system.

ref_gen Reference genome version
library_name Name of the reference library

Value

Loci-based DNA-mutational fingerprint of the cancer cell line as found in the input VCF file.

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```
parse_vcf_query_into_db
```

parse_vcf_query_into_db This function adds the variants of parsed custom CCLs to a monet DB instance

Description

parse_vcf_query_into_db This function adds the variants of parsed custom CCLs to a monet DB instance

Usage

```
parse_vcf_query_into_db(
   g_query,
   ref_gen = "GRCH37",
   library_name,
   test_mode = FALSE
)
```

Arguments

g_query a GenomicRanges object

ref_gen a character string specifying the reference genome version. All training sets are

associated with a reference genome version. Default is "GRCH37".

library_name a character string giving the name of the library to add the cancer cell lines to.

Default is "CUSTOM". Library name will be automatically added as a suffix to

the identifier.

test_mode Is this a test? Just for internal use

Value

Message wheather the adding was successful

Description

This function procides information on the reference library names

```
read_library_names(ref_gen)
```

Arguments

ref_gen

a character vector specifying the reference genome version. All training sets are associated with a reference genome version. Default is "GRCH37".

Value

Returns a character vector of the contained libraries

Examples

```
read_library_names(ref_gen = "GRCH37")
```

Description

Read the GRange object for a specific library

Usage

```
read_mutation_grange_objects(
   library_name,
   mutational_weight_inclusion_threshold,
   ref_gen,
   test_mode
)
```

Arguments

```
library_name a character string giving the name of the library
```

 ${\tt mutational_weight_inclusion_threshold}$

a numerical giving the lower bound for mutational weight to be included

ref_gen Reference genome version. All training sets are associated with a reference

genome version. Default: GRCH37

test_mode Is this a test? Just for internal use

Value

The R Table sim_list which contains the CoSMIC CLP fingerprints

```
remove_ccls_from_database
```

Remove Cancer Cell Line

Description

This function removes a cancer cell line training fingerprint (VCF file) from the database. The names of all training sets can be seen by using the function show_contained_cls.

Usage

Arguments

A character vector giving the names of the cancer cell line identifiers to be removed. Can be one or many

ref_gen A character vector specifying the reference genome version. All training sets

are associated with a reference genome version. Default is "GRCH37".

library_name Name of the library from which ccls are to be removed

test_mode Signifies if this is a test run

Value

Message that indicates whether the removal was succesful.

Examples

```
remove_ccls_from_database(
    ccl_names = "HT29",
    ref_gen = "GRCH37",
    library_name = "CELLMINER",
    test_mode = TRUE
)
```

remove_library_from_database

Remove entire Library from Database

Description

This function removes a entire library from the database by removing all associated cancer cell line fingerprints from the database.

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Usage

```
remove_library_from_database(library, ref_gen = "GRCH37", test_mode = FALSE)
```

Arguments

library a character vector giving the names of the library to be removed.

ref_gen a character vector specifying the reference genome version. All training sets are

associated with a reference genome version. Default is "GRCH37".

test_mode is this a test? Just for internal use.

Value

Message that indicates whether the removal was succesful.

Examples

show_contained_ccls show contained ccls

Description

This function displays the names, amount of mutations and the overall weight of the mutations of all contained cancer cell line fingerprints for a chosen reference genome and optional library.

Usage

```
show_contained_ccls(ref_gen, verbose)
```

Arguments

ref_gen a character vector specifying the reference genome version. All training sets are

associated with a reference genome version. Default is "GRCH37".

verbose Should DB informations be printed?

Value

R table which contains identifiers of all cancer cell line samples which match the specified parameters (reference genome and library).

```
## Show all contained cancer cell lines for reference GRCH37:
show_contained_ccls(ref_gen = "GRCH37", verbose = TRUE)
```

```
show_contained_variants_for_ccl

Variants In Cancer Cell Line
```

This function shows all mutations present in the database for a selected cancer cell line and reference genome.

Usage

```
show_contained_variants_for_ccl(
   name_ccl,
   ref_gen,
   library_name,
   mutational_weight_inclusion_threshold)
```

Arguments

name_ccl a character vector giving the identifier of the cancer cell line for which mutations will be shown.

ref_gen a character vector specifying the reference genome version. All training sets are

associated with a reference genome version. Default is "GRCH37".

library_name Name of the reference library mutational_weight_inclusion_threshold

Include only mutations with a weight of at least x. Range: 0.0 to 1.0. 1= unique to CL. $\sim 0 =$ found in many CCL samples.

Value

GenomicRanges object that contains the ccl's variants

```
## Show all mutations for Cancer Cell Line 'SK_OV_3'
show_contained_variants_for_ccl(
   name_ccl = "SK_OV_3",
   ref_gen = "GRCH37",
   library_name = "CELLMINER",
   mutational_weight_inclusion_threshold = 0
)
```

```
show_contained_variants_in_library

All variants contained in reference library
```

This function shows all variants contained in a reference library for a given inclusion weight. Default inclusion weight is 0 (all variants).

Usage

```
show_contained_variants_in_library(
    ref_gen,
    library_name,
    mutational_weight_inclusion_threshold)
```

Arguments

Value

Returns a GenomicRanges object that contains the variants

```
## Show all variants contained in reference library CELLMINER
show_contained_variants_in_library(
    ref_gen = "GRCH37",
    library_name = "CELLMINER",
    mutational_weight_inclusion_threshold = 0
)
```

```
show_which_ccls_contain_variant
```

Cancer cell lines with specific variant

Description

This function displays all cancer cell lines in the database which contain a specified variant. Utilizes closed interval coordinates.

Usage

```
show_which_ccls_contain_variant(
    start,
    end,
    chromosome,
    ref_gen,
    library_name,
    mutational_weight_inclusion_threshold)
```

Arguments

start Start coordinate end Stop coordinate

chromosome, 'chr' prefixes are ignored

ref_gen a character vector specifying the reference genome version. All training sets are

associated with a reference genome version. Default is "GRCH37".

library_name Name of the reference library mutational_weight_inclusion_threshold

Include only mutations with a weight of at least x. Range: 0.0 to 1.0. 1= unique

to CL. ~ 0 = found in many CCL samples.

Value

Returns a GenomicRanges object that contains the variant if present. Member ccls can be found in the \$Member_ccl vector

```
show_which_ccls_contain_variant(
    start = 92030762,
    end = 92030762,
    chromosome = 8,
    ref_gen = "GRCH37",
    library_name = "CELLMINER",
    mutational_weight_inclusion_threshold = 0
)
```

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