Package 'Neoantimon'

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Title Neoantimon: A multifunctional R package for identification of tumor-specific neoantigens

Type Package

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Description This Package has been developed to calculate candidates neoantigens from Mutation Data (.vcf,.txt,string).
License MIT + file LICENSE
VignetteBuilder knitr
Encoding UTF-8
Depends R (>= 3.3.0)
biocViews
Imports ensemblVEP, devtools, graphics, grDevices, stats, utils, biomaRt Suggests data.table, knitr, rmarkdown LazyData FALSE RoxygenNote 7.1.0 R topics documented:
CalculatePriorityScores 2 Export_Summary_Entire_Fragments 3 Export_Summary_Fragments 4 Export_Summary_IndelSV 5 Export_Summary_IndelSV_perFragments 6 Export_Summary_SNV 7 MainEntireRegionClass1 8 MainINDELClass2 10 MainINDELClass1 12 MainSeqFragmentClass2 15 MainSeqFragmentClass2 21 MainSNVClass1 23 MainSNVClass2 27

Index		44
	TestAnalysis	43
	sample_vcf.vep	
	sample_vcf.snps	
	sample_vcf.annovar	
	sample_sv_bnd	
	sample_rna_exp	
	sample_result_SVFusion_CLASS2_ALL	
	sample_result_SVFusion_CLASS1_ALL	
	sample_result_SNV_CLASS2_ALL	
	sample_result_SNV_CLASS1_ALL	
	sample_result_SeqFragment_CLASS2_ALL	
	sample_result_SeqFragment_CLASS1_ALL	
	sample_result_INDEL_CLASS2_ALL	
	sample_result_INDEL_CLASS1_ALL	
	sample_refMrna.grch37.fa	
	sample_refFlat.grch37	
	sample_hla_table_c2	
	sample_hla_table_c1	
	sample_copynum	
	MainSVFUSIONClass2	34
	MainSVFUSIONClass1	31

 ${\tt CalculatePriorityScores}$

Calculate priority scores

Description

Calculate priority scores

Usage

CalculatePriorityScores(result, useRNAvaf = FALSE)

Arguments

result A results file generated from main functions

useRNAvaf To indicate whether this function uses DNA VAF or RNA VAF for the calcula-

tion.

Value

P_I: Priority score using the IC50.

P_R: Priority score using the percentage of rank affinity.

P: Priority score implemented in MuPeXI (Bjerregaard et al. 2017).

```
{\it Export\_Summary\_Entire\_Fragments} \\ {\it Export\_Summary\_Count\ from\ Indel/SV\ Results}
```

Description

Export Summary Count from Indel/SV Results

Usage

```
Export_Summary_Entire_Fragments(
   Input,
   Mut_IC50_th = NA,
   Mut_Rank_th = NA,
   Total_RNA_th = NA,
   Tumor_RNA_th = NA,
   MutRatio_th = NA,
   WriteLongIndel = NA,
   DupCount = FALSE
)
```

Arguments

Input	Input file generated from MainSNVClass1,2.
Mut_IC50_th	The threshold for mutant peptide to be neoantigen by IC50.
Mut_Rank_th	The threshold for mutant peptide to be neoantigen by Rank.
Total_RNA_th	The total RNA expression threshold.
Tumor_RNA_th	The tumor specific RNA expression threshold.
MutRatio_th	The mutation ratio threshold.
WriteLongIndel	If setting a file name, Write Long Indels of which the p-value is less than 0.05 .
DupCount	Count for each different HLA type

Value

Num_Alteration The number of evaluated alterations.

Num_Alteration_Generating_NeoAg The number of evaluated alterations that can generate neoantigen.

Num_Peptide The number of evaluated peptifdes.

```
Export_Summary_Fragments
```

Export Summary Count from Indel/SV Results

Description

Export Summary Count from Indel/SV Results

Usage

```
Export_Summary_Fragments(
   Input,
   Mut_IC50_th = NA,
   Mut_Rank_th = NA,
   Total_RNA_th = NA,
   Tumor_RNA_th = NA,
   MutRatio_th = NA,
   WriteLongIndel = NA,
   DupCount = FALSE
)
```

Arguments

Input	Input file generated from MainSNVClass1,2.
Mut_IC50_th	The threshold for mutant peptide to be neoantigen by IC50.
Mut_Rank_th	The threshold for mutant peptide to be neoantigen by Rank.
Total_RNA_th	The total RNA expression threshold.
Tumor_RNA_th	The tumor specific RNA expression threshold.
MutRatio_th	The mutation ratio threshold.
WriteLongIndel	If setting a file name, Write Long Indels of which the p-value is less than 0.05.
DupCount	Count for each different HLA type

Value

Num_Alteration The number of evaluated alterations.

Num_Alteration_Generating_NeoAg The number of evaluated alterations that can generate neoantigen.

Num_Peptide The number of evaluated peptifdes.

```
Export_Summary_IndelSV
```

Export Summary Count from Indel/SV Results

Description

Export Summary Count from Indel/SV Results

Usage

```
Export_Summary_IndelSV(
   Input,
   Mut_IC50_th = NA,
   Mut_Rank_th = NA,
   Total_RNA_th = NA,
   Tumor_RNA_th = NA,
   MutRatio_th = NA,
   Weight = NA,
   WriteLongIndel = NA,
   IgnoreLongIndel = 0,
   DupCount = FALSE
)
```

Arguments

Input	Input file generated from MainSNVClass1,2.
Mut_IC50_th	The threshold for mutant peptide to be neoantigen by IC50.
Mut_Rank_th	The threshold for mutant peptide to be neoantigen by Rank.
Total_RNA_th	The total RNA expression threshold.
Tumor_RNA_th	The tumor specific RNA expression threshold.
MutRatio_th	The mutation ratio threshold.
Weight	The weight for alterations.
WriteLongIndel	If setting a file name, Write Long Indels of which the p-value is less than 0.05.
IgnoreLongIndel	
	Ignore Indels of which p-value is less than the indicated value for counting.
DupCount	Count for each different HLA type

Value

Num_Alteration The number of evaluated alterations.

Num_Alteration_Generating_NeoAg The number of evaluated alterations that can generate neoantigen.

Num_Peptide The number of evaluated peptifdes.

```
Export_Summary_IndelSV_perFragments
```

Export Summary Count from Indel/SV Results

Description

Export Summary Count from Indel/SV Results

Usage

```
Export_Summary_IndelSV_perFragments(
   Input,
   Mut_IC50_th = NA,
   Mut_Rank_th = NA,
   Total_RNA_th = NA,
   Tumor_RNA_th = NA,
   MutRatio_th = NA,
   Weight = NA,
   WriteLongIndel = NA,
   IgnoreLongIndel = 0,
   DupCount = FALSE
)
```

Arguments

Input	Input file generated from MainSNVClass1,2.
Mut_IC50_th	The threshold for mutant peptide to be neoantigen by IC50.
Mut_Rank_th	The threshold for mutant peptide to be neoantigen by Rank.
Total_RNA_th	The total RNA expression threshold.
Tumor_RNA_th	The tumor specific RNA expression threshold.
MutRatio_th	The mutation ratio threshold.
Weight	The weight for alterations.
WriteLongIndel	If setting a file name, Write Long Indels of which the p-value is less than 0.05.
IgnoreLongIndel	
	Ignore Indels of which p-value is less than the indicated value for counting.
DupCount	Count for each different HLA type

Value

Num_Alteration The number of evaluated alterations.

Num_Alteration_Generating_NeoAg The number of evaluated alterations that can generate neoantigen.

Num_Peptide The number of evaluated peptifdes.

Export_Summary_SNV

Export Summary Count from SNV Results

Description

Export Summary Count from SNV Results

Usage

```
Export_Summary_SNV(
   Input,
   Mut_IC50_th = NA,
   Mut_Rank_th = NA,
   Wt_IC50_th = NA,
   Wt_Rank_th = NA,
   Total_RNA_th = NA,
   Tumor_RNA_th = NA,
   MutRatio_th = NA,
   DupCount = FALSE
)
```

Arguments

Input	Input file generated from MainSNVClass1,2.
Mut_IC50_th	The threshold for mutant peptide to be neoantigen.
Mut_Rank_th	The threshold for mutant peptide to be neoantigen.
Wt_IC50_th	The threshold for wt peptide to be neoantigen.
Wt_Rank_th	The threshold for wt peptide to be neoantigen.
Total_RNA_th	The total RNA expression threshold.
Tumor_RNA_th	The tumor specific RNA expression threshold.
MutRatio_th	The mutation ratio threshold.
DupCount	Count for each different HLA type

Value

Num_Alteration The number of evaluated alterations.

Num_Alteration_Generating_NeoAg The number of evaluated alterations that can generate neoantigen.

Num_Peptide The number of evaluated peptifdes.

MainEntireRegionClass1

Calculate A Set All Neoantigen Candidates from A Given Gene Symbol and nm_id for MHC Class1 (Not yet stably available)

Description

Calculate A Set All Neoantigen Candidates from A Given Gene Symbol and nm_id for MHC Class1 (Not yet stably available)

Usage

```
MainEntireRegionClass1(
  input_nm_id,
 group_ids = seq(1:length(input_nm_id)),
 hla_file = "here_is_a_table",
 hla_{types} = NA,
 file_name_in_hla_table = NA,
 refflat_file = paste(hmdir, "lib/refFlat.txt", sep = "/"),
  refmrna_file = paste(hmdir, "lib/refMrna.fa", sep = "/"),
 hmdir = getwd(),
  job_id = "ID",
 export_dir = paste("result", job_id, "EntireRegion1", sep = "."),
 netMHCpan_dir = paste(hmdir, "lib/netMHCpan-4.0/netMHCpan", sep = "/"),
 peptide_length = c(8, 9, 10, 11, 12, 13),
 reading_frame = 1,
 CalculateIC50 = FALSE,
  ignore_short = TRUE
)
```

Arguments

input_nm_id	(Required) An input amino acid sequence indicated as NM_ID
group_ids	flag to cluster the same group
hla_file	A tab separated file indicating HLA types. The 1st column is input_file name, and the following columns indicate HLA types.
	See by data(sample_hla_table_c1); sample_hla_table_c1;
hla_types	Set a list of HLA types
file_name_in_hl	a_table
	If the name (1st column) in HLA table is not the same as input_file, indicate the corresponding name (Default=input_file).
refflat_file	refFlat file to be used in constructing peptide. (Default=paste(hmdir, "lib/refFlat.txt", sep="").
	See "https://github.com/hase62/Neoantimon"
refmrna_file	refMrna file to be used in constructing peptide (Default=paste(hmdir, "lib/refMrna.fa", sep="").
	See "https://github.com/hase62/Neoantimon"
hmdir	Home directory for the analysis (Default = getwd()).

job_id Job-id to be attached in output files (Default = "NO_job_id").

export_dir The directory will be stored results (Default = "paste("result", file_name_in_hla_table,

job_id, sep=".")")

netMHCpan_dir The file directory to netMHCpan (Default="lib/netMHCpan-4.0/netMHCpan").

peptide_length Peptide Length to be generated (Default = 8.9.10.11.12.13). reading_frame The starting frame of the input sequence (Default = 1)

 ${\tt CalculateIC50} \quad Whether \ Calculate \ IC50 \ by \ NetMHC pan \ or \ not.$

ignore_short Ignore to output results of Short Peptide Less Than min(peptide_length)

Value

void (Calculated Neoantigen Files will be generated as .tsv files.):

HLA: HLA type used to calculate neoantigen.

Pos: The position of a fraction of peptide used to be evaluated from the full-length peptide.

Gene: Gene symbol used to be evaluated in NetMHCpan.

Evaluated_Mutant_Peptide: The mutant peptide to be evaluated.

Evaluated_Mutant_Peptide_Core: The core peptide of the mutant peptide to be evaluated in NetMHC-

pan.

Mut_IC50: IC50 value for evaluated mutant peptide.

Mut_Rank: Rank value for evaluated mutanat peptide.

Chr: Chromosome Number of the mutation.

NM_ID: NM_ID used to construct peptides from the mutation.

Change: The annotation to be described in .vcf file.

Ref: reference type nucleic acid base.

Alt: alternative type nucleic acid base.

Prob: A probability of reference nucleic acid base described in .vcf file.

Mutation_Prob: A probability of alternative nucleic acid base described in .vcf file.

Exon_Start: The exon start position of the corrsponding NM_ID.

Exon_End: The exon end position of the corrsponding NM_ID.

Mutation_Position: The mutation position of the corrsponding NM_ID.

Total_Depth: The sum depth of the reference and alternative nucleic acid base.

Tumor_Depth: The depth of the alternative nucleic acid base.

Wt_Peptide: The full-length of the wild-type peptide.

Mutant_Peptide: The full-length of the mutant peptide.

Total_RNA: The expression amount of the corresponding RNA.

Tumor_RNA_Ratio: The variant allele frequency of the corresponding RNA.

Tumor_RNA: The modified amount of the corresponding RNA level based on RNA Reads.

Tumor_RNA_based_on_DNA: The modified amount of the corresponding RNA level based on DNA Reads.

MutRatio: The mean value of the cancer cell fraction probability.

MutRatio_Min: The 1% percentile of the cancer cell fraction probability.

MutRatio_Max: The 99% percentile of the cancer cell fraction probability.

MainEntireRegionClass2

Calculate A Set All Neoantigen Candidates from A Given Gene Symbol and nm_id for MHC Class2 (Not yet stably available)

Description

Calculate A Set All Neoantigen Candidates from A Given Gene Symbol and nm_id for MHC Class2 (Not yet stably available)

Usage

```
MainEntireRegionClass2(
  input_nm_id,
 group_ids = seq(1:length(input_nm_id)),
 hla_file = "here_is_a_table",
 hla_{types} = NA,
 file_name_in_hla_table = NA,
 refflat_file = paste(hmdir, "lib/refFlat.txt", sep = "/"),
  refmrna_file = paste(hmdir, "lib/refMrna.fa", sep = "/"),
 hmdir = getwd(),
  job_id = "ID",
 export_dir = paste("result", job_id, "EntireRegion2", sep = "."),
 netMHCIIpan_dir = paste(hmdir, "lib/netMHCIIpan-3.1/netMHCIIpan", sep = "/"),
 peptide_length = c(15),
 reading_frame = 1,
 CalculateIC50 = FALSE,
 ignore_short = TRUE
)
```

Arguments

input_nm_id	(Required) An input amino acid sequence indicated as NM_ID
group_ids	flag to cluster the same group
hla_file	A tab separated file indicating HLA types. The 1st column is input_file name, and the following columns indicate HLA types.
	See by data(sample_hla_table_c1); sample_hla_table_c1;
hla_types	Set a list of HLA types
file_name_in_hla_table	
	If the name (1st column) in HLA table is not the same as input_file, indicate the corresponding name (Default=input_file).
refflat_file	refFlat file to be used in constructing peptide. (Default=paste(hmdir, "lib/refFlat.txt", sep="").
	See "https://github.com/hase62/Neoantimon"
refmrna_file	refMrna file to be used in constructing peptide (Default=paste(hmdir, "lib/refMrna.fa", sep="").
	See "https://github.com/hase62/Neoantimon"
hmdir	Home directory for the analysis (Default = getwd()).

job_id Job-id to be attached in output files (Default = "NO_job_id").

export_dir The directory will be stored results (Default = "paste("result", file_name_in_hla_table,

job_id, sep=".")")

netMHCIIpan_dir

The file directory to netMHCpan (Default="lib/netMHCIIpan-3.2/netMHCIIpan").

$$\label{eq:peptide_length} \begin{split} & \text{peptide_length} & \text{Peptide Length to be generated (Default} = 8,9,10,11,12,13). \\ & \text{reading_frame} & \text{The starting frame of the input sequence (Default} = 1) \end{split}$$

 ${\tt CalculateIC50} \quad Whether \ Calculate \ IC50 \ by \ NetMHC pan \ or \ not.$

ignore_short Ignore to output results of Short Peptide Less Than min(peptide_length)

Value

void (Calculated Neoantigen Files will be generated as .tsv files.):

HLA: HLA type used to calculate neoantigen.

Pos: The position of a fraction of peptide used to be evaluated from the full-length peptide.

Gene: Gene symbol used to be evaluated in NetMHCpan.

Evaluated_Mutant_Peptide: The mutant peptide to be evaluated.

Evaluated_Mutant_Peptide_Core: The core peptide of the mutant peptide to be evaluated in NetMHC-pan.

Mut_IC50: IC50 value for evaluated mutant peptide.

Mut_Rank: Rank value for evaluated mutanat peptide.

Chr: Chromosome Number of the mutation.

NM_ID: NM_ID used to construct peptides from the mutation.

Change: The annotation to be described in .vcf file.

Ref: reference type nucleic acid base.

Alt: alternative type nucleic acid base.

Prob: A probability of reference nucleic acid base described in .vcf file.

Mutation Prob: A probability of alternative nucleic acid base described in .vcf file.

Exon_Start: The exon start position of the corrsponding NM_ID.

Exon_End: The exon end position of the corrsponding NM_ID.

Mutation_Position: The mutation position of the corrsponding NM_ID.

Total_Depth: The sum depth of the reference and alternative nucleic acid base.

Tumor_Depth: The depth of the alternative nucleic acid base.

Wt_Peptide: The full-length of the wild-type peptide.

Mutant_Peptide: The full-length of the mutant peptide.

Total RNA: The expression amount of the corresponding RNA.

Tumor_RNA_Ratio: The variant allele frequency of the corresponding RNA.

Tumor RNA: The modified amount of the corresponding RNA level based on RNA Reads.

Tumor_RNA_based_on_DNA: The modified amount of the corresponding RNA level based on DNA Reads.

MutRatio: The mean value of the cancer cell fraction probability.

MutRatio_Min: The 1% percentile of the cancer cell fraction probability.

MutRatio_Max: The 99% percentile of the cancer cell fraction probability.

MainINDELClass1

Calculate Neoantigen Candidates on INDELs for MHC Class1

Description

Calculate Neoantigen Candidates on INDELs for MHC Class1

Usage

```
MainINDELClass1(
  input_annovar_format_file = NA,
  input_vep_format_file = NA,
  input_vcf_format_file_and_vep = NA,
  hla_file = "here_is_a_table",
  hla_{types} = NA,
  file_name_in_hla_table = "sample",
  refflat_file = paste(hmdir, "lib/refFlat.txt", sep = "/"),
  refmrna_file = paste(hmdir, "lib/refMrna.fa", sep = "/"),
  hmdir = getwd(),
  job_id = "ID",
  export_dir = paste("result", job_id, "INDEL1", sep = "."),
  rnaexp_file = NA,
  rnabam_file = NA,
  cnv_file = NA,
  purity = 1,
  netMHCpan_dir = paste(hmdir, "lib/netMHCpan-4.0/netMHCpan", sep = "/"),
  MHCflurry = NA,
  refdna_file = NA,
  samtools_dir = "samtools",
  bcftools_dir = NA,
  chr_column = NA,
  mutation_start_column = NA,
  mutation_end_column = NA,
  mutation_ref_column = NA,
  mutation_alt_column = NA,
  nm_id_column = NA,
  depth_normal_column = NA,
  depth_tumor_column = NA,
  ambiguous_between_exon = 0,
  ambiguous_codon = 0,
  peptide_length = c(8, 9, 10, 11, 12, 13),
  ignore_short = TRUE,
  SNPs = NA,
  multiple_variants = FALSE
)
```

Arguments

```
input_annovar_format_file
```

An input vcf file annotated by ANNOVAR (http://annovar.openbioinformatics.org/en/latest/). You can directly indicate a matrix, which is the same as annovar format vcf file, as input.

See by data(sample_vcf.annovar); sample_vcf.annovar.txt;

input_vep_format_file

An input file annotated by Ensembl Variant Effect Predictor (VEP). You can directly indicate a matrix, which is the same as annovar format VEP file, as input.

See by data(sample_vcf.vep); sample_vcf.vep.txt;

input_vcf_format_file_and_vep

An input vcf file and path to Ensembl Variant Effect Predictor (VEP). Before us-

ing this option, please install vep according to the official cite ("https://asia.ensembl.org/info/docs/tool

13

hla_file A tab separated file indicating HLA types. The 1st column is input_file name,

and the following columns indicate HLA types.

See by data(sample_hla_table_c1); sample_hla_table_c1;

hla_types Set a list of HLA types

file_name_in_hla_table

If the name (1st column) in HLA table is not the same as input_file, indicate the

corresponding name.

refflat_file refFlat file to be used in constructing peptide. (Default=paste(hmdir, "lib/refFlat.txt",

sep="").

See "https://github.com/hase62/Neoantimon"

refmrna_file refMrna file to be used in constructing peptide (Default=paste(hmdir, "lib/refMrna.fa",

sep="").

See "https://github.com/hase62/Neoantimon"

hmdir Home directory for the analysis (Default = getwd()).

job_id Job-id to be attached in output files (Default = "NO_job_id").

export_dir The directory will be stored results (Default = "paste("result", file_name_in_hla_table,

job_id, sep=".")")

rnaexp_file A file including RNA expressions (Default=NA). The 1st, 2nd and 3rd columns

are "GeneSymbol Chr:Exonstart-Exonend (locus) ExpressionAmount", respec-

tively. The 1st row should be any header.

See by data(sample_rna_exp); sample_rna_exp;

rnabam_file RNA bam file to calculate variant allele frequency of RNA at each mutation

(Default=NA).

cnv_file A file including copy number variation to calculate cancer cell fraction prob-

ability (CCFP) (Default=NA). The format is according to ASCAT output files. The columns are "SNPName Chromosome Position LogR segmentedLogR BAF segmentedBAF CopyNumber MinorAllele RawCopyNumber" The 1st row should

be the above header.

See data(sample_copynum); sample_copynum;

purity Tumor purity or tumor contents ratio required to calculate CCFP (Default=1).

netMHCpan_dir The file directory to netMHCpan (Default="lib/netMHCpan-4.0/netMHCpan").

MHCflurry Output MHCflurry results. Return a list of both results (Default=FALSE).

refdna_file refdna_file information to be used to calculate RNA VAF (Default=NA).

See "https://github.com/hase62/Neoantimon"

samtools_dir The file directory to samtools_0_x_x (Default="samtools"). It shouled be indi-

cated when you indicate RNA-bam and try to calculate RNA VAF.

bcftools_dir The file directory to netMHCpan (Default="bcftools"). It shouled be indicated when you indicate RNA-bam and try to calculate RNA VAF. samtools 0_x_x includes bcftools in the directory.

chr_column The column number describing chromosome number in input_file (Default=NA, but will automatically search "Chr" in header).

mutation_start_column

The column number describing mutation start Position in input_file (Default=NA, but will automatically search "Start" in header).

mutation_end_column

The column number describing mutation end Position in input_file (Default=NA, but will automatically search "End" in header).

mutation_ref_column

The column number describing mutation Ref in input_file (Default=NA, but will automatically search "Ref" in header).

mutation_alt_column

The column number describing mutation Alt in input_file (Default=NA, but will automatically search "Alt" in header).

nm_id_column The column number describing NM IDs in input_file such as

"SLCO1C1:NM_001145944:exon7:c.692_693insG:p.L231fs" (Default=NA).

depth_normal_column

The column number describing the read count from normal cells (Default = NA).

depth_tumor_column

 $\label{eq:continuous} The \ column \ number \ describing \ the \ read \ count \ from \ tumor \ cells \ (Default=NA).$ $\ ambiguous_between_exon$

The maximum number to permit the differences between Exon-Lengths from refFlat and refMrna (Default=0).

ambiguous_codon

The maximum number to permit the differences between inputfile- and refMrnaoriented translation start/end position (Default=0).

peptide_length Peptide Length to be generated (Default = 8,9,10,11,12,13).

ignore_short Ignore to output results of short peptide less than min (peptide_length)

SNPs Apply indivisual SNPs on peptides by indicate a vcf file.

multiple_variants

Reflect multiple variants on a peptide, e.g., SNVs on frameshift region.

Value

void (Calculated Neoantigen Files will be generated as .tsv files.):

HLA: HLA type used to calculate neoantigen.

Pos: The position of a fraction of peptide used to be evaluated from the full-length peptide.

Gene Gene symbol used to be evaluated in NetMHCpan.

Evaluated_Mutant_Peptide: The mutant peptide to be evaluated.

Evaluated_Mutant_Peptide_Core: The core peptide of the mutant peptide to be evaluated in NetMHC-pan.

Mut_IC50: IC50 value for evaluated mutant peptide.

Mut_Rank: Rank value for evaluated mutanat peptide.

Chr: Chromosome Number of the mutation.

NM_ID: NM_ID used to construct peptides from the mutation.

Change: The annotation to be described in .vcf file.

Ref: reference type nucleic acid base.

Alt: alternative type nucleic acid base.

Prob: A probability of reference nucleic acid base described in .vcf file.

Mutation_Prob: A probability of alternative nucleic acid base described in .vcf file.

Exon_Start: The exon start position of the corrsponding NM_ID.

Exon_End: The exon end position of the corrsponding NM_ID.

Mutation_Position: The mutation position of the corrsponding NM_ID.

Total_Depth: The sum depth of the reference and alternative nucleic acid base.

Tumor_Depth: The depth of the alternative nucleic acid base.

Wt_Peptide: The full-length of the wild-type peptide.

Mutant_Peptide: The full-length of the mutant peptide.

Total_RNA: The expression amount of the corresponding RNA.

Tumor_RNA_Ratio: The variant allele frequency of the corresponding RNA.

Tumor_RNA: The modified amount of the corresponding RNA level based on RNA Reads.

Tumor_RNA_based_on_DNA: The modified amount of the corresponding RNA level based on DNA Reads.

MutRatio: The mean value of the cancer cell fraction probability.

MutRatio_Min: The 1% percentile of the cancer cell fraction probability.

MutRatio_Max: The 99% percentile of the cancer cell fraction probability.

P_I: Priority score using the IC50. Please use CalculatePriorityScores <- function(result, useR-NAvaf = FALSE)

P_R: Priority score using the percentage of rank affinity. Please use CalculatePriorityScores <-function(result, useRNAvaf = FALSE)

P: Priority score implemented in MuPeXI (Bjerregaard et al. 2017). Please use CalculatePriorityScores <- function(result, useRNAvaf = FALSE)

MainINDELClass2

Calculate Neoantigen Candidates on INDELs for MHC Class2

Description

Calculate Neoantigen Candidates on INDELs for MHC Class2

Usage

```
MainINDELClass2(
  input_annovar_format_file = NA,
  input_vep_format_file = NA,
  input_vcf_format_file_and_vep = NA,
  hla_file = "here_is_a_table",
  hla_types = NA,
  file_name_in_hla_table = "sample",
```

```
refflat_file = paste(hmdir, "lib/refFlat.txt", sep = "/"),
      refmrna_file = paste(hmdir, "lib/refMrna.fa", sep = "/"),
      hmdir = getwd(),
      job_id = "ID",
      export_dir = paste("result", job_id, "INDEL2", sep = "."),
      rnaexp_file = NA,
      rnabam_file = NA,
      cnv_file = NA,
      purity = 1,
      netMHCIIpan_dir = paste(hmdir, "lib/netMHCIIpan-3.1/netMHCIIpan", sep = "/"),
      refdna_file = NA,
      samtools_dir = "samtools",
      bcftools_dir = NA,
      chr_column = NA,
      mutation_start_column = NA,
      mutation_end_column = NA,
      mutation_ref_column = NA,
      mutation_alt_column = NA,
      nm_id_column = NA,
      depth_normal_column = NA,
      depth_tumor_column = NA,
      ambiguous_between_exon = 0,
      ambiguous_codon = 0,
      peptide_length = c(15),
      ignore_short = TRUE,
      SNPs = NA,
      multiple_variants = FALSE
    )
Arguments
    input_annovar_format_file
                     An input vcf file annotated by ANNOVAR (http://annovar.openbioinformatics.org/en/latest/).
                     You can directly indicate a matrix, which is the same as annovar format vcf file,
                     as input.
                     See by data(sample_vcf.annovar); sample_vcf.annovar.txt;
    input_vep_format_file
                     An input file annotated by Ensembl Variant Effect Predictor (VEP). You can
                     directly indicate a matrix, which is the same as annovar format VEP file, as
                     See by data(sample_vcf.vep); sample_vcf.vep.txt;
    input_vcf_format_file_and_vep
                     An input vcf file and path to Ensembl Variant Effect Predictor (VEP). Before us-
                     ing this option, please install vep according to the official cite ("https://asia.ensembl.org/info/docs/tool
   hla_file
                     A tab separated file indicating HLA types. The 1st column is input_file name,
                     and the following columns indicate HLA types.
                     See by data(sample_hla_table_c1); sample_hla_table_c1;
    hla_types
                     Set a list of HLA types
    file_name_in_hla_table
                     If the name (1st column) in HLA table is not the same as input_file, indicate the
```

corresponding name.

refflat file refFlat file to be used in constructing peptide. (Default=paste(hmdir, "lib/refFlat.txt", sep=""). See "https://github.com/hase62/Neoantimon" refmrna_file refMrna file to be used in constructing peptide (Default=paste(hmdir, "lib/refMrna.fa", sep=""). See "https://github.com/hase62/Neoantimon" hmdir Home directory for the analysis (Default = getwd()). job_id Job-Id to be attached in output files (Default = "NO_job_id"). export_dir The directory will be stored results (Default = "paste("result", file_name_in_hla_table, job_id, sep=".")") A file including RNA expressions (Default=NA). The 1st, 2nd and 3rd columns rnaexp_file are "GeneSymbol Chr:Exonstart-Exonend (locus) ExpressionAmount", respectively. The 1st row should be any header. See by data(sample_rna_exp); sample_rna_exp; rnabam_file RNA bam file to calculate variant allele frequency of RNA at each mutation (Default=NA). cnv_file A file including copy number variation to calculate cancer cell fraction probability (CCFP) (Default=NA). The format is according to ASCAT output files. The columns are "SNPName Chromosome Position LogR segmentedLogR BAF segmentedBAF CopyNumber MinorAllele RawCopyNumber" The 1st row should be the above header. See data(sample_copynum); sample_copynum; Tumor purity or tumor contents ratio required to calculate CCFP (Default=1). purity netMHCIIpan_dir The file directory to netMHCpan (Default="lib/netMHCIIpan-3.2/netMHCpan"). refdna file information to be used to calculate RNA VAF (Default=NA). refdna_file See "https://github.com/hase62/Neoantimon" The file directory to samtools 0 x x (Default="samtools"). It shouled be indisamtools dir cated when you indicate RNA-bam and try to calculate RNA VAF. bcftools dir The file directory to netMHCpan (Default="bcftools"). It shouled be indicated when you indicate RNA-bam and try to calculate RNA VAF. samtools 0_x_x

chr_column

The column number describing chromosome number in input_file (Default=NA, but will automatically search "Chr" in header).

includes beftools in the directory.

mutation_start_column

The column number describing mutation start Position in input_file (Default=NA, but will automatically search "Start" in header).

mutation_end_column

The column number describing mutation end Position in input_file (Default=NA, but will automatically search "End" in header).

mutation_ref_column

The column number describing mutation Ref in input_file (Default=NA, but will automatically search "Ref" in header).

mutation_alt_column

The column number describing mutation Alt in input_file (Default=NA, but will automatically search "Alt" in header).

nm_id_column The column number describing NM IDs in input_file such as

"SLCO1C1:NM_001145944:exon7:c.692_693insG:p.L231fs" (Default=NA).

depth_normal_column

The column number describing the read count from normal cells (Default = NA).

depth_tumor_column

The column number describing the read count from tumor cells (Default = NA).

ambiguous_between_exon

The maximum number to permit the differences between Exon-Lengths from

refFlat and refMrna (Default=0).

ambiguous_codon

The maximum number to permit the differences between inputfile- and refMrna-

oriented translation start/end position (Default=0).

peptide_length Peptide Length to be generated (Default = 15 in HLA Class2).

ignore_short Ignore to output results of short peptide less than min (peptide_length)

SNPs Apply indivisual SNPs on peptides by indicate a vcf file.

multiple_variants

Reflect multiple variants on a peptide, e.g., SNVs on frameshift region.

Value

void (Calculated Neoantigen Files will be generated as .tsv files.):

HLA: HLA type used to calculate neoantigen.

Pos: The position of a fraction of peptide used to be evaluated from the full-length peptide.

Gene Gene symbol used to be evaluated in NetMHCpan.

Evaluated_Mutant_Peptide: The mutant peptide to be evaluated.

Evaluated_Mutant_Peptide_Core: The core peptide of the mutant peptide to be evaluated in NetMHC-pan.

Mut_IC50: IC50 value for evaluated mutant peptide.

Mut Rank: Rank value for evaluated mutanat peptide.

Chr: Chromosome Number of the mutation.

NM ID: NM ID used to construct peptides from the mutation.

Change: The annotation to be described in .vcf file.

Ref: reference type nucleic acid base.

Alt: alternative type nucleic acid base.

Prob: A probability of reference nucleic acid base described in .vcf file.

Mutation_Prob: A probability of alternative nucleic acid base described in .vcf file.

Exon_Start: The exon start position of the corrsponding NM_ID.

Exon_End: The exon end position of the corrsponding NM_ID.

Mutation_Position: The mutation position of the corrsponding NM_ID.

Total_Depth: The sum depth of the reference and alternative nucleic acid base.

Tumor_Depth: The depth of the alternative nucleic acid base.

Wt_Peptide: The full-length of the wild-type peptide.

Mutant_Peptide: The full-length of the mutant peptide.

Total_RNA: The expression amount of the corresponding RNA.

Tumor_RNA_Ratio: The variant allele frequency of the corresponding RNA.

Tumor_RNA: The modified amount of the corresponding RNA level based on RNA Reads.

Tumor_RNA_based_on_DNA: The modified amount of the corresponding RNA level based on DNA Reads.

MutRatio: The mean value of the cancer cell fraction probability.

MutRatio_Min: The 1% percentile of the cancer cell fraction probability.

MutRatio_Max: The 99% percentile of the cancer cell fraction probability.

P_I: Priority score using the IC50. Please use CalculatePriorityScores <- function(result, useR-NAvaf = FALSE)

P_R: Priority score using the percentage of rank affinity. Please use CalculatePriorityScores <-function(result, useRNAvaf = FALSE)

P: Priority score implemented in MuPeXI (Bjerregaard et al. 2017). Please use CalculatePriorityScores <- function(result, useRNAvaf = FALSE)

 $\begin{tabular}{ll} Main Seq Fragment Class 1 & Calculate Neoantigen Candidates from A Given Sequence for MHC \\ & Class 1 \end{tabular}$

Description

Calculate Neoantigen Candidates from A Given Sequence for MHC Class1

Usage

```
MainSeqFragmentClass1(
  input_sequence = NA,
  group_ids = seq(1:length(reference_nm_id)),
  hla_file = "here_is_a_table",
  hla_{types} = NA,
  file_name_in_hla_table = NA,
  refflat_file = paste(hmdir, "lib/refFlat.txt", sep = "/"),
  refmrna_file = paste(hmdir, "lib/refMrna.fa", sep = "/"),
  hmdir = getwd(),
  job_id = "ID",
  export_dir = paste("result", job_id, "SeqFragment1", sep = "."),
  netMHCpan_dir = paste(hmdir, "lib/netMHCpan-4.0/netMHCpan", sep = "/"),
  peptide_length = c(8, 9, 10, 11, 12, 13),
  reference_nm_id = NA,
  reference_gene_symbol = NA,
  ignore_short = TRUE
)
```

Arguments

```
input_sequence (Required) An input amino acid sequence
group_ids flag to cluster the same group
hla_file A tab separated file indicating HLA types. The 1st c
```

A tab separated file indicating HLA types. The 1st column is input_file name,

and the following columns indicate HLA types.

See by data(sample_hla_table_c1); sample_hla_table_c1;

hla_types Set a list of HLA types

file_name_in_hla_table

If the name (1st column) in HLA table is not the same as input_file, indicate the

corresponding name (Default=input_file).

refflat_file refFlat file to be used in constructing peptide. (Default=paste(hmdir, "lib/refFlat.txt",

sep="").

See "https://github.com/hase62/Neoantimon"

refmrna_file refMrna file to be used in constructing peptide (Default=paste(hmdir, "lib/refMrna.fa",

sep="").

See "https://github.com/hase62/Neoantimon"

hmdir Home directory for the analysis (Default = getwd()).

job_id Job-Id to be attached in output files (Default = "NO_job_id").

export_dir The directory will be stored results (Default = "paste("result", file_name_in_hla_table,

job_id, sep=".")")

netMHCpan_dir The file directory to netMHCpan (Default="lib/netMHCpan-4.0/netMHCpan").

peptide_length Peptide Length to be generated (Default = 8,9,10,11,12,13).

reference_nm_id

Corresponding original sequences that the input sequence is generated. If franctions of peptides generated from the input are included in the indicated protein, such peptides are removed. It can be indicated when gene_symbol is not NA.

reference_gene_symbol

Corresponding original sequences that the input sequence is generated. If franctions of peptides generated from the input are included in the indicated protein, such peptides are removed. It can be indicated when nm_id is not NA.

ignore_short Ignore to output results of short peptide less than min (peptide_length)

Value

void (Calculated Neoantigen Files will be generated as .tsv files.):

HLA: HLA type used to calculate neoantigen.

Pos: The position of a fraction of peptide used to be evaluated from the full-length peptide.

Gene Gene symbol used to be evaluated in NetMHCpan.

Evaluated_Mutant_Peptide: The mutant peptide to be evaluated.

Evaluated_Mutant_Peptide_Core: The core peptide of the mutant peptide to be evaluated in NetMHC-pan.

Mut_IC50: IC50 value for evaluated mutant peptide.

Mut_Rank: Rank value for evaluated mutanat peptide.

Chr: Chromosome Number of the mutation.

NM_ID: NM_ID used to construct peptides from the mutation.

Change: The annotation to be described in .vcf file.

Ref: reference type nucleic acid base.

Alt: alternative type nucleic acid base.

Prob: A probability of reference nucleic acid base described in .vcf file.

Mutation_Prob: A probability of alternative nucleic acid base described in .vcf file.

Exon_Start: The exon start position of the corrsponding NM_ID.

Exon_End: The exon end position of the corrsponding NM_ID.

Mutation_Position: The mutation position of the corrsponding NM_ID.

Total_Depth: The sum depth of the reference and alternative nucleic acid base.

Tumor_Depth: The depth of the alternative nucleic acid base.

Wt_Peptide: The full-length of the wild-type peptide.

Mutant_Peptide: The full-length of the mutant peptide.

Total_RNA: The expression amount of the corresponding RNA.

Tumor_RNA_Ratio: The variant allele frequency of the corresponding RNA.

Tumor RNA: The modified amount of the corresponding RNA level based on RNA Reads.

Tumor_RNA_based_on_DNA: The modified amount of the corresponding RNA level based on DNA Reads.

MutRatio: The mean value of the cancer cell fraction probability.

MutRatio_Min: The 1% percentile of the cancer cell fraction probability.

MutRatio_Max: The 99% percentile of the cancer cell fraction probability.

P_I: Priority score using the IC50. Please use CalculatePriorityScores <- function(result, useR-NAvaf = FALSE)

P_R: Priority score using the percentage of rank affinity. Please use CalculatePriorityScores <- function(result, useRNAvaf = FALSE)

P: Priority score implemented in MuPeXI (Bjerregaard et al. 2017). Please use CalculatePriorityScores <- function(result, useRNAvaf = FALSE)

 $\begin{tabular}{lll} Main Seq Fragment Class 2 & Calculate Neoantigen Candidates from A Given Sequence for MHC \\ & Class 2 \\ \end{tabular}$

Description

Calculate Neoantigen Candidates from A Given Sequence for MHC Class2

Usage

```
MainSeqFragmentClass2(
   input_sequence = NA,
   group_ids = seq(1:length(reference_nm_id)),
   hla_file = "here_is_a_table",
   hla_types = NA,
   file_name_in_hla_table = NA,
   refflat_file = paste(hmdir, "lib/refFlat.txt", sep = "/"),
   refmrna_file = paste(hmdir, "lib/refMrna.fa", sep = "/"),
   hmdir = getwd(),
   job_id = "ID",
   export_dir = paste("result", job_id, "SeqFragment2", sep = "."),
   netMHCIIpan_dir = paste(hmdir, "lib/netMHCIIpan-3.1/netMHCIIpan", sep = "/"),
   peptide_length = c(15),
   reference_nm_id = NA,
```

```
reference_gene_symbol = NA,
ignore_short = TRUE
)
```

Arguments

input_sequence (Required) An input amino acid sequence

group_ids flag to cluster the same group

hla_file A tab separated file indicating HLA types. The 1st column is input_file name,

and the following columns indicate HLA types.

See by data(sample_hla_table_c1); sample_hla_table_c1;

hla_types Set a list of HLA types

file_name_in_hla_table

If the name (1st column) in HLA table is not the same as input_file, indicate the

corresponding name (Default=input_file).

refflat_file refFlat file to be used in constructing peptide. (Default=paste(hmdir, "lib/refFlat.txt",

sep="").

See "https://github.com/hase62/Neoantimon"

refmrna_file refMrna file to be used in constructing peptide (Default=paste(hmdir, "lib/refMrna.fa",

sep="").

See "https://github.com/hase62/Neoantimon"

hmdir Home directory for the analysis (Default = getwd()).

job_id Job-Id to be attached in output files (Default = "NO_job_id").

export_dir The directory will be stored results (Default = "paste("result", file_name_in_hla_table,

job_id, sep=".")")

netMHCIIpan_dir

The file directory to netMHCpan (Default="lib/netMHCIIpan-3.2/netMHCIIpan").

peptide_length Peptide Length to be generated (Default = 8,9,10,11,12,13).

reference_nm_id

Corresponding original sequences that the input sequence is generated. If franctions of peptides generated from the input are included in the indicated protein, such peptides are removed. It can be indicated when gene_symbol is not NA.

reference_gene_symbol

Corresponding original sequences that the input sequence is generated. If franctions of peptides generated from the input are included in the indicated protein,

such peptides are removed. It can be indicated when nm_id is not NA.

ignore_short Ignore to output results of short peptide less than min (peptide_length)

Value

void (Calculated Neoantigen Files will be generated as .tsv files.):

HLA: HLA type used to calculate neoantigen.

Pos: The position of a fraction of peptide used to be evaluated from the full-length peptide.

Gene Gene symbol used to be evaluated in NetMHCpan.

Evaluated_Mutant_Peptide: The mutant peptide to be evaluated.

Evaluated_Mutant_Peptide_Core: The core peptide of the mutant peptide to be evaluated in NetMHC-pan.

MainSNVClass1 23

Mut_IC50: IC50 value for evaluated mutant peptide.

Mut_Rank: Rank value for evaluated mutanat peptide.

Chr: Chromosome Number of the mutation.

NM_ID: NM_ID used to construct peptides from the mutation.

Change: The annotation to be described in .vcf file.

Ref: reference type nucleic acid base.

Alt: alternative type nucleic acid base.

Prob: A probability of reference nucleic acid base described in .vcf file.

Mutation_Prob: A probability of alternative nucleic acid base described in .vcf file.

Exon_Start: The exon start position of the corrsponding NM_ID.

Exon_End: The exon end position of the corrsponding NM_ID.

Mutation_Position: The mutation position of the corrsponding NM_ID.

Total_Depth: The sum depth of the reference and alternative nucleic acid base.

Tumor Depth: The depth of the alternative nucleic acid base.

Wt_Peptide: The full-length of the wild-type peptide.

Mutant_Peptide: The full-length of the mutant peptide.

Total_RNA: The expression amount of the corresponding RNA.

Tumor_RNA_Ratio: The variant allele frequency of the corresponding RNA.

Tumor_RNA: The modified amount of the corresponding RNA level based on RNA Reads.

Tumor_RNA_based_on_DNA: The modified amount of the corresponding RNA level based on DNA Reads.

MutRatio: The mean value of the cancer cell fraction probability.

MutRatio_Min: The 1% percentile of the cancer cell fraction probability.

MutRatio_Max: The 99% percentile of the cancer cell fraction probability.

P_I: Priority score using the IC50. Please use CalculatePriorityScores <- function(result, useR-NAvaf = FALSE)

P_R: Priority score using the percentage of rank affinity. Please use CalculatePriorityScores <-function(result, useRNAvaf = FALSE)

P: Priority score implemented in MuPeXI (Bjerregaard et al. 2017). Please use CalculatePriorityScores <- function(result, useRNAvaf = FALSE)

MainSNVClass1

Calculate Neoantigen Candidates on SNVs for MHC Class1

Description

Calculate Neoantigen Candidates on SNVs for MHC Class1

24 MainSNVClass1

Usage

```
MainSNVClass1(
      input_annovar_format_file = NA,
      input_vep_format_file = NA,
      input_vcf_format_file_and_vep = NA,
     hla_file = "here_is_a_table",
     hla_{types} = NA,
      file_name_in_hla_table = "sample",
      refflat_file = paste(hmdir, "lib/refFlat.txt", sep = "/"),
      refmrna_file = paste(hmdir, "lib/refMrna.fa", sep = "/"),
     hmdir = getwd(),
      job_id = "ID",
      export_dir = paste("result", job_id, "SNV1", sep = "."),
      rnaexp_file = NA,
      rnabam_file = NA,
      cnv_file = NA,
     purity = 1,
     netMHCpan_dir = paste(hmdir, "lib/netMHCpan-4.0/netMHCpan", sep = "/"),
     MHCflurry = NA,
     refdna_file = NA,
      samtools_dir = "samtools",
     bcftools_dir = NA,
      chr_column = NA,
     mutation_start_column = NA,
     mutation_end_column = NA,
     mutation_ref_column = NA,
     mutation_alt_column = NA,
     nm_id_column = NA,
     depth_normal_column = NA,
      depth_tumor_column = NA,
      ambiguous_between_exon = 0,
      ambiguous_codon = 0,
      peptide_length = c(8, 9, 10, 11, 12, 13),
      ignore_short = TRUE,
      SNPs = NA,
     multiple_variants = FALSE
   )
Arguments
    input_annovar_format_file
                    An input vcf file annotated by ANNOVAR (http://annovar.openbioinformatics.org/en/latest/).
                    You can directly indicate a matrix, which is the same as annovar format vcf file,
                    as input.
                    See by data(sample_vcf.annovar); sample_vcf.annovar.txt;
    input_vep_format_file
                    An input file annotated by Ensembl Variant Effect Predictor (VEP). You can
                    directly indicate a matrix, which is the same as annovar format VEP file, as
                    See by data(sample_vcf.vep); sample_vcf.vep.txt;
    input_vcf_format_file_and_vep
```

A list of (1) An input vcf file, (2) path to Ensembl Variant Effect Predictor

MainSNVClass1 25

(VEP), and (3) cache file for VEP. Before using this option, please install vep according to the official cite ("https://asia.ensembl.org/info/docs/tools/vep/index.html"). hla_file A tab separated file indicating HLA types. The 1st column is input_file name, and the following columns indicate HLA types. See by data(sample_hla_table_c1); sample_hla_table_c1; Set a list of HLA types hla_types file_name_in_hla_table If the name (1st column) in HLA table is not the same as input_file, indicate the corresponding name. refflat_file refFlat file to be used in constructing peptide. (Default=paste(hmdir, "lib/refFlat.txt", See "https://github.com/hase62/Neoantimon" refMrna file to be used in constructing peptide (Default=paste(hmdir, "lib/refMrna.fa", refmrna_file sep=""). See "https://github.com/hase62/Neoantimon" hmdir Home directory for the analysis (Default = getwd()). Job-id to be attached in output files (Default = "NO job id"). job_id export_dir The directory will be stored results (Default = "paste("result", file_name_in_hla_table, job id, sep=".")") A file including RNA expressions (Default=NA). The 1st, 2nd and 3rd columns rnaexp_file are "GeneSymbol Chr:Exonstart-Exonend (locus) ExpressionAmount", respectively. The 1st row should be any header. See by data(sample_rna_exp); sample_rna_exp; RNA bam file to calculate variant allele frequency of RNA at each mutation rnabam_file (Default=NA). cnv_file A file including copy number variation to calculate cancer cell fraction probability (CCFP) (Default=NA). The format is according to ASCAT output files. The columns are "SNPName Chromosome Position LogR segmentedLogR BAF segmentedBAF CopyNumber MinorAllele RawCopyNumber" The 1st row should be the above header. See data(sample_copynum); sample_copynum; Tumor purity or tumor contents ratio required to calculate CCFP (Default=1). purity netMHCpan_dir The file directory to netMHCpan (Default="lib/netMHCpan-4.0/netMHCpan"). Output MHCflurry results. Return a list of both results (Default=FALSE). MHCflurry refdna file refdna_file information to be used to calculate RNA VAF (Default=NA). See "https://github.com/hase62/Neoantimon" The file directory to samtools_0_x_x (Default="samtools"). It shouled be indisamtools_dir cated when you indicate RNA-bam and try to calculate RNA VAF. The file directory to netMHCpan (Default="bcftools"). It shouled be indicated bcftools_dir when you indicate RNA-bam and try to calculate RNA VAF . samtools 0_x_x includes beftools in the directory. The column number describing chromosome number in input_file (Default=NA, chr_column but will automatically search "Chr" in header). mutation_start_column

The column number describing mutation start Position in input_file (Default=NA,

but will automatically search "Start" in header).

26 MainSNVClass1

mutation_end_column

The column number describing mutation end Position in input_file (Default=NA, but will automatically search "End" in header).

mutation_ref_column

The column number describing mutation Ref in input_file (Default=NA, but will automatically search "Ref" in header).

mutation_alt_column

The column number describing mutation Alt in input_file (Default=NA, but will automatically search "Alt" in header).

nm_id_column The column number describing NM IDs in input_file such as

"SLCO1C1:NM_001145944:exon7:c.692_693insG:p.L231fs" (Default=NA).

depth_normal_column

The column number describing the read count from normal cells (Default = NA).

depth_tumor_column

The column number describing the read count from tumor cells (Default = NA).

ambiguous_between_exon

The maximum number to permit the differences between Exon-Lengths from refFlat and refMrna (Default=0).

ambiguous_codon

The maximum number to permit the differences between inputfile- and refMrnaoriented translation start/end position (Default=0).

peptide_length Peptide Length to be generated (Default = 8,9,10,11,12,13).

ignore_short Ignore to output results of short peptide less than min (peptide_length)

SNPs Apply indivisual SNPs on peptides by indicate a vcf file.

multiple_variants

Reflect multiple variants on a peptide, e.g., SNVs on frameshift region.

Value

void (Calculated Neoantigen Files will be generated as .tsv files.)

HLA: HLA type used to calculate neoantigen.

Pos: The position of a fraction of peptide used to be evaluated from the full-length peptide.

Gene Gene symbol used to be evaluated in NetMHCpan.

Evaluated_Mutant_Peptide: The mutant peptide to be evaluated.

Mut_IC50: IC50 value for evaluated mutant peptide.

Mut_Rank: Rank value for evaluated mutanat peptide.

Evaluated_Wt_Peptide: The wild-type peptide to be evaluated.

Wt_IC50: IC50 value for evaluated wild-type peptide.

Wt_Rank: Rank value for evaluated wild-type peptide.

Chr: Chromosome Number of the mutation.

NM_ID: NM_ID used to construct peptides from the mutation.

Change: The annotation to be described in .vcf file.

Ref: reference type nucleic acid base.

Alt: alternative type nucleic acid base.

Prob: A probability of reference nucleic acid base described in .vcf file.

MainSNVClass2 27

Mutation_Prob: A probability of alternative nucleic acid base described in .vcf file.

Exon_Start: The exon start position of the corrsponding NM_ID.

Exon_End: The exon end position of the corrsponding NM_ID.

Mutation_Position: The mutation position of the corrsponding NM_ID.

Total_Depth: The sum depth of the reference and alternative nucleic acid base.

Tumor_Depth: The depth of the alternative nucleic acid base.

Wt_Peptide: The full-length of the wild-type peptide.

Mutant_Peptide: The full-length of the mutant peptide.

Total_RNA: The expression amount of the corresponding RNA.

Tumor_RNA_Ratio: The variant allele frequency of the corresponding RNA.

Tumor_RNA: The modified amount of the corresponding RNA level based on RNA Reads.

Tumor_RNA_based_on_DNA: The modified amount of the corresponding RNA level based on DNA Reads.

MutRatio: The mean value of the cancer cell fraction probability.

MutRatio_Min: The 1% percentile of the cancer cell fraction probability.

MutRatio_Max: The 99% percentile of the cancer cell fraction probability.

P_I: Priority score using the IC50. Please use CalculatePriorityScores <- function(result, useR-NAvaf = FALSE)

P_R: Priority score using the percentage of rank affinity. Please use CalculatePriorityScores <-function(result, useRNAvaf = FALSE)

P: Priority score implemented in MuPeXI (Bjerregaard et al. 2017). Please use CalculatePriorityScores <- function(result, useRNAvaf = FALSE)

MainSNVClass2

Calculate Neoantigen Candidates on SNVs for MHC Class2

Description

Calculate Neoantigen Candidates on SNVs for MHC Class2

Usage

```
MainSNVClass2(
  input_annovar_format_file = NA,
  input_vep_format_file = NA,
  input_vcf_format_file_and_vep = NA,
  hla_file = "here_is_a_table",
  hla_types = NA,
  file_name_in_hla_table = "sample",
  refflat_file = paste(hmdir, "lib/refFlat.txt", sep = "/"),
  refmrna_file = paste(hmdir, "lib/refMrna.fa", sep = "/"),
  hmdir = getwd(),
  job_id = "ID",
  export_dir = paste("result", job_id, "SNV2", sep = "."),
  rnaexp_file = NA,
```

28 MainSNVClass2

rnabam_file = NA,

```
cnv file = NA.
      purity = 1,
      netMHCIIpan_dir = paste(hmdir, "lib/netMHCIIpan-3.2/netMHCIIpan", sep = "/"),
      refdna_file = NA,
      samtools_dir = "samtools",
      bcftools_dir = NA,
      chr_column = NA,
      mutation_start_column = NA,
      mutation_end_column = NA,
      mutation_ref_column = NA,
      mutation_alt_column = NA,
      nm_id_column = NA,
      depth_normal_column = NA,
      depth_tumor_column = NA,
      ambiguous_between_exon = 0,
      ambiguous_codon = 0,
      peptide_length = c(15),
      ignore_short = TRUE,
      SNPs = NA,
      multiple_variants = FALSE
    )
Arguments
    input_annovar_format_file
                      An input vcf file annotated by ANNOVAR (http://annovar.openbioinformatics.org/en/latest/).
                      You can directly indicate a matrix, which is the same as annovar format vcf file,
                      as input. See by data(sample_vcf.annovar); sample_vcf.annovar.txt;
    input_vep_format_file
                      An input file annotated by Ensembl Variant Effect Predictor (VEP). You can
                      directly indicate a matrix, which is the same as annovar format VEP file, as
                      input.
                      See by data(sample_vcf.vep); sample_vcf.vep.txt;
    input_vcf_format_file_and_vep
                      A list of (1) An input vcf file, (2) path to Ensembl Variant Effect Predictor
                      (VEP), and (3) cache file for VEP. Before using this option, please install vep ac-
                      cording to the official cite ("https://asia.ensembl.org/info/docs/tools/vep/index.html").
    hla_file
                      A tab separated file indicating HLA types. The 1st column is input_file name,
                      and the following columns indicate HLA types.
                      See by data(sample_hla_table_c1); sample_hla_table_c1;
                      Set a list of HLA types
    hla_types
    file_name_in_hla_table
                      If the name (1st column) in HLA table is not the same as input_file, indicate the
                      corresponding name.
                      refFlat file to be used in constructing peptide. (Default=paste(hmdir, "lib/refFlat.txt",
    refflat_file
                      sep="").
                      See "https://github.com/hase62/Neoantimon"
    refmrna_file
                      refMrna file to be used in constructing peptide (Default=paste(hmdir, "lib/refMrna.fa",
                      See "https://github.com/hase62/Neoantimon"
```

MainSNVClass2 29

Home directory for the analysis (Default = getwd()).

hmdir

job_id Job-Id to be attached in output files (Default = "NO_job_id"). The directory will be stored results (Default = "paste("result", file_name_in_hla_table, export_dir job_id, sep=".")") rnaexp_file A file including RNA expressions (Default=NA). The 1st, 2nd and 3rd columns are "GeneSymbol Chr:Exonstart-Exonend (locus) ExpressionAmount", respectively. The 1st row should be any header. See by data(sample_rna_exp); sample_rna_exp; RNA bam file to calculate variant allele frequency of RNA at each mutation rnabam_file (Default=NA). cnv_file A file including copy number variation to calculate cancer cell fraction probability (CCFP) (Default=NA). The format is according to ASCAT output files. The columns are "SNPName Chromosome Position LogR segmentedLogR BAF segmentedBAF CopyNumber MinorAllele RawCopyNumber" The 1st row should be the above header. See data(sample_copynum); sample_copynum; purity Tumor purity or tumor contents ratio required to calculate CCFP (Default=1). netMHCIIpan_dir The file directory to netMHCpan (Default="lib/netMHCIIpan-3.2/netMHCpan"). refdna_file refdna_file information to be used to calculate RNA VAF (Default=NA). See "https://github.com/hase62/Neoantimon" samtools_dir The file directory to samtools_0_x_x (Default="samtools"). It shouled be indicated when you indicate RNA-bam and try to calculate RNA VAF. The file directory to netMHCpan (Default="bcftools"). It shouled be indicated bcftools_dir when you indicate RNA-bam and try to calculate RNA VAF . samtools 0 x x includes beftools in the directory. chr_column The column number describing chromosome number in input_file (Default=NA, but will automatically search "Chr" in header). mutation_start_column The column number describing mutation start Position in input_file (Default=NA, but will automatically search "Start" in header). mutation_end_column The column number describing mutation end Position in input_file (Default=NA, but will automatically search "End" in header). mutation_ref_column The column number describing mutation Ref in input file (Default=NA, but will automatically search "Ref" in header). mutation_alt_column The column number describing mutation Alt in input_file (Default=NA, but will automatically search "Alt" in header). nm_id_column The column number describing NM IDs in input_file such as "SLCO1C1:NM_001145944:exon7:c.692_693insG:p.L231fs" (Default=NA). depth_normal_column The column number describing the read count from normal cells (Default = NA). depth_tumor_column

The column number describing the read count from tumor cells (Default = NA).

30 MainSNVClass2

ambiguous_between_exon

The maximum number to permit the differences between Exon-Lengths from refFlat and refMrna (Default=0).

ambiguous_codon

The maximum number to permit the differences between inputfile- and refMrna-

oriented translation start/end position (Default=0).

peptide_length Peptide Length to be generated (Default = 15 in HLA Class2).

ignore_short Ignore to output results of short peptide less than min (peptide_length)

SNPs Apply indivisual SNPs on peptides by indicate a vcf file.

multiple_variants

Reflect multiple variants on a peptide, e.g., SNVs on frameshift region.

Value

void (Calculated Neoantigen Files will be generated as .tsv files.):

HLA: HLA type used to calculate neoantigen.

Pos: The position of a fraction of peptide used to be evaluated from the full-length peptide.

Gene Gene symbol used to be evaluated in NetMHCpan.

Evaluated_Mutant_Peptide: The mutant peptide to be evaluated.

Mut_IC50: IC50 value for evaluated mutant peptide.

Mut_Rank: Rank value for evaluated mutanat peptide.

Evaluated_Wt_Peptide: The wild-type peptide to be evaluated.

Wt_IC50: IC50 value for evaluated wild-type peptide.

Wt_Rank: Rank value for evaluated wild-type peptide.

Chr: Chromosome Number of the mutation.

NM_ID: NM_ID used to construct peptides from the mutation.

Change: The annotation to be described in .vcf file.

Ref: reference type nucleic acid base.

Alt: alternative type nucleic acid base.

Prob: A probability of reference nucleic acid base described in .vcf file.

Mutation_Prob: A probability of alternative nucleic acid base described in .vcf file.

Exon_Start: The exon start position of the corrsponding NM_ID.

Exon_End: The exon end position of the corrsponding NM_ID.

Mutation Position: The mutation position of the corrsponding NM ID.

Total_Depth: The sum depth of the reference and alternative nucleic acid base.

Tumor_Depth: The depth of the alternative nucleic acid base.

Wt_Peptide: The full-length of the wild-type peptide.

Mutant_Peptide: The full-length of the mutant peptide.

Total_RNA: The expression amount of the corresponding RNA.

Tumor_RNA_Ratio: The variant allele frequency of the corresponding RNA.

Tumor_RNA: The modified amount of the corresponding RNA level based on RNA Reads.

Tumor_RNA_based_on_DNA: The modified amount of the corresponding RNA level based on DNA Reads.

MainSVFUSIONClass1 31

MutRatio: The mean value of the cancer cell fraction probability.

MutRatio_Min: The 1% percentile of the cancer cell fraction probability.

MutRatio_Max: The 99% percentile of the cancer cell fraction probability.

P_I: Priority score using the IC50. Please use CalculatePriorityScores <- function(result, useR-NAvaf = FALSE)

P_R: Priority score using the percentage of rank affinity. Please use CalculatePriorityScores <-function(result, useRNAvaf = FALSE)

P: Priority score implemented in MuPeXI (Bjerregaard et al. 2017). Please use CalculatePriorityScores <- function(result, useRNAvaf = FALSE)

MainSVFUSIONClass1

Calculate Neoantigen Candidates on SV fusions for MHC Class1

Description

Calculate Neoantigen Candidates on SV fusions for MHC Class1

Usage

```
MainSVFUSIONClass1(
  input_file,
  hla_file = "here_is_a_table",
  hla_{types} = NA,
  file_name_in_hla_table = input_file,
  refflat_file = paste(hmdir, "lib/refFlat.txt", sep = "/"),
  refmrna_file = paste(hmdir, "lib/refMrna.fa", sep = "/"),
  hmdir = getwd(),
  job_id = "ID",
  export_dir = paste("result", job_id, "SV1", sep = "."),
  rnaexp_file = NA,
  rnabam_file = NA,
  cnv_file = NA,
  purity = 1,
  netMHCpan_dir = paste(hmdir, "lib/netMHCpan-4.0/netMHCpan", sep = "/"),
  refdna_file = NA,
  samtools_dir = NA,
  bcftools_dir = NA,
  chr_column = NA,
  mutation_start_column = NA,
  mutation_end_column = NA,
  mutation_ref_column = NA,
  mutation_alt_bnd_column = NA,
  depth_normal_column = NA,
  depth_tumor_column = NA,
  nm_id_column = NA,
  ambiguous_between_exon = 0,
  ambiguous_codon = 0,
  peptide_length = c(8, 9, 10, 11, 12, 13),
  gene_symbol_column = NA,
```

32 MainSVFUSIONClass1

```
mate_id_column = NA,
ignore_short = TRUE
)
```

Arguments

input_file (Required) An input vcf file (BND format) annotated by,

e.g., ANNOVAR (http://annovar.openbioinformatics.org/en/latest/) or other soft-

wares.

See by data(sample_sv_bnd); sample_sv_bnd;

hla_file A tab separated file indicating HLA types. The 1st column is input_file name,

and the following columns indicate HLA types.

See by data(sample_hla_table_c1); sample_hla_table_c1;

hla_types Set a list of HLA types

file_name_in_hla_table

If the name (1st column) in HLA table is not the same as input_file, indicate the

corresponding name (Default=input_file).

refflat_file refFlat file to be used in constructing peptide. (Default=paste(hmdir, "lib/refFlat.txt",sep="").

See "https://github.com/hase62/Neoantimon"

refmrna_file refMrna file to be used in constructing peptide (Default=paste(hmdir, "lib/refMrna.fa",

sep="").

See "https://github.com/hase62/Neoantimon"

hmdir Home directory for the analysis (Default = getwd()).

job_id Job-Id to be attached in output files (Default = "NO job id").

export_dir The directory will be stored results (Default = "paste("result", file_name_in_hla_table,

job_id, sep=".")")

rnaexp_file A file including RNA expressions (Default=NA). The 1st, 2nd and 3rd columns

are "GeneSymbol Chr:Exonstart-Exonend (locus) ExpressionAmount", respec-

tively. The 1st row should be any header.

See by data(sample_rna_exp); sample_rna_exp;

rnabam_file RNA bam file to calculate variant allele frequency of RNA at each mutation

(Default=NA).

cnv_file A file including copy number variation to calculate cancer cell fraction prob-

ability (CCFP) (Default=NA). The format is according to ASCAT output files. The columns are "SNPName Chromosome Position LogR segmentedLogR BAF segmentedBAF CopyNumber MinorAllele RawCopyNumber" The 1st row should

be the above header.

See data(sample_copynum); sample_copynum;

purity Tumor purity or tumor contents ratio required to calculate CCFP (Default=1).

netMHCpan_dir The file directory to netMHCpan (Default="lib/netMHCpan-4.0/netMHCpan").

refdna_file refdna_file information to be used to calculate RNA VAF (Default=NA).

See "https://github.com/hase62/Neoantimon"

samtools_dir The file directory to samtools_0_x_x (Default="samtools"). It shouled be indi-

cated when you indicate RNA-bam and try to calculate RNA VAF.

bcftools_dir The file directory to netMHCpan (Default="bcftools"). It shouled be indicated

when you indicate RNA-bam and try to calculate RNA VAF. samtools 0_x_x

includes beftools in the directory.

MainSVFUSIONClass1 33

chr_column The column number describing chromosome number in input_file (Default=NA, but will automatically search "Chr" in header).

mutation_start_column

The column number describing mutation start Position in input_file (Default=NA, but will automatically search "Start" in header) .

mutation_end_column

The column number describing mutation end Position in input_file (Default=NA, but will automatically search "End" in header).

mutation_ref_column

The column number describing mutation Ref in input_file (Default=NA, but will automatically search "Ref" in header).

mutation_alt_bnd_column

The column number describing mutation Alt (BND format) in input_file (Default=NA, but will automatically search "Alt" in header).

depth_normal_column

The column number describing the read count from normal cells (Default = NA).

depth_tumor_column

The column number describing the read count from tumor cells (Default = NA).

nm_id_column (Required if gene_symbol_column = NA) The column number describing NM IDs in input_file such as

"SLCO1C1:NM 001145944:exon7:c.692 693insG:p.L231fs" (Default=NA).

ambiguous_between_exon

The maximum number to permit the differences between Exon-Lengths from refFlat and refMrna (Default=0).

ambiguous_codon

The maximum number to permit the differences between inputfile- and refMrnaoriented translation start/end position (Default=0).

peptide_length Peptide Length to be generated (Default = 8,9,10,11,12,13).

gene_symbol_column

(Required if nm_id_column = NA) The column number describing gene symbol in input_file (Default=NA).

mate_id_column (Required) The column indicating mateIDs or svIDs such as "SVMERGE1_1" (Default=NA).

ignore_short Ignore to output results of short peptide less than min (peptide_length)

Value

void (Calculated Neoantigen Files will be generated as .tsv files.):

HLA: HLA type used to calculate neoantigen.

Pos: The position of a fraction of peptide used to be evaluated from the full-length peptide.

Gene Gene symbol used to be evaluated in NetMHCpan.

Evaluated_Mutant_Peptide: The mutant peptide to be evaluated.

Evaluated_Mutant_Peptide_Core: The core peptide of the mutant peptide to be evaluated in NetMHC-pan.

Mut_IC50: IC50 value for evaluated mutant peptide.

Mut_Rank: Rank value for evaluated mutanat peptide.

Chr: Chromosome Number of the mutation.

34 MainSVFUSIONClass2

NM_ID: NM_ID used to construct peptides from the mutation.

Change: The annotation to be described in .vcf file.

Ref: reference type nucleic acid base.

Alt: alternative type nucleic acid base.

Prob: A probability of reference nucleic acid base described in .vcf file.

Mutation_Prob: A probability of alternative nucleic acid base described in .vcf file.

Exon_Start: The exon start position of the corrsponding NM_ID.

Exon_End: The exon end position of the corrsponding NM_ID.

Mutation_Position: The mutation position of the corrsponding NM_ID.

Total_Depth: The sum depth of the reference and alternative nucleic acid base.

Tumor_Depth: The depth of the alternative nucleic acid base.

Wt_Peptide: The full-length of the wild-type peptide.

Mutant_Peptide: The full-length of the mutant peptide.

Total_RNA: The expression amount of the corresponding RNA.

Tumor_RNA_Ratio: The variant allele frequency of the corresponding RNA.

Tumor_RNA: The modified amount of the corresponding RNA level based on RNA Reads.

Tumor_RNA_based_on_DNA: The modified amount of the corresponding RNA level based on DNA Reads.

MutRatio: The mean value of the cancer cell fraction probability.

MutRatio_Min: The 1% percentile of the cancer cell fraction probability.

MutRatio_Max: The 99% percentile of the cancer cell fraction probability.

P_I: Priority score using the IC50. Please use CalculatePriorityScores <- function(result, useR-NAvaf = FALSE)

P_R: Priority score using the percentage of rank affinity. Please use CalculatePriorityScores <-function(result, useRNAvaf = FALSE)

P: Priority score implemented in MuPeXI (Bjerregaard et al. 2017). Please use CalculatePriorityScores <- function(result, useRNAvaf = FALSE)

MainSVFUSIONClass2

Calculate Neoantigen Candidates on SV fusions for MHC Class2

Description

Calculate Neoantigen Candidates on SV fusions for MHC Class2

Usage

```
MainSVFUSIONClass2(
  input_file,
  hla_file = "here_is_a_table",
  hla_types = NA,
  file_name_in_hla_table = input_file,
  refflat_file = paste(hmdir, "lib/refFlat.txt", sep = "/"),
  refmrna_file = paste(hmdir, "lib/refMrna.fa", sep = "/"),
```

MainSVFUSIONClass2 35

```
hmdir = getwd(),
  job_id = "ID",
  export_dir = paste("result", job_id, "SV2", sep = "."),
  rnaexp_file = NA,
  rnabam_file = NA,
  cnv_file = NA,
 purity = 1,
  netMHCIIpan_dir = paste(hmdir, "lib/netMHCIIpan-3.1/netMHCIIpan", sep = "/"),
  refdna_file = NA,
  samtools_dir = NA,
 bcftools_dir = NA,
 chr_column = NA,
 mutation_start_column = NA,
 mutation_end_column = NA,
 mutation_ref_column = NA,
 mutation_alt_bnd_column = NA,
 depth_normal_column = NA,
  depth_tumor_column = NA,
 nm_id_column = NA,
  ambiguous_between_exon = 0,
  ambiguous_codon = 0,
 peptide_length = c(15),
 gene_symbol_column = NA,
 mate_id_column = NA,
  ignore_short = TRUE
)
```

Arguments

export_dir

job_id, sep=".")")

input_file (Required) An input vcf file (BND format) annotated by, e.g., ANNOVAR (http://annovar.openbioinformatics.org/en/latest/) or other softwares. See by data(sample_sv_bnd); sample_sv_bnd; A tab separated file indicating HLA types. The 1st column is input_file name, hla_file and the following columns indicate HLA types. See by data(sample_hla_table_c1); sample_hla_table_c1; Set a list of HLA types hla_types file_name_in_hla_table If the name (1st column) in HLA table is not the same as input_file, indicate the corresponding name (Default=input_file). refFlat file to be used in constructing peptide. (Default=paste(hmdir, "lib/refFlat.txt",sep=""). refflat_file See "https://github.com/hase62/Neoantimon" refMrna file to be used in constructing peptide (Default=paste(hmdir, "lib/refMrna.fa", refmrna_file sep=""). See "https://github.com/hase62/Neoantimon" hmdir Home directory for the analysis (Default = getwd()). job_id Job-Id to be attached in output files (Default = "NO_job_id").

The directory will be stored results (Default = "paste("result", file_name_in_hla_table,

36 MainSVFUSIONClass2

rnaexp_file A file including RNA expressions (Default=NA). The 1st, 2nd and 3rd columns are "GeneSymbol Chr:Exonstart-Exonend (locus) ExpressionAmount", respectively. The 1st row should be any header.

See by data(sample_rna_exp); sample_rna_exp;

rnabam_file RNA bam file to calculate variant allele frequency of RNA at each mutation (Default=NA).

A file including copy number variation to calculate cancer cell fraction probability (CCFP) (Default=NA). The format is according to ASCAT output files. The columns are "SNPName Chromosome Position LogR segmentedLogR BAF segmentedBAF CopyNumber MinorAllele RawCopyNumber" The 1st row should be the above header.

See data(sample_copynum); sample_copynum;

purity Tumor purity or tumor contents ratio required to calculate CCFP (Default=1). netMHCIIpan_dir

The file directory to netMHCpan (Default="lib/netMHCIIpan-3.2/netMHCpan").

refdna_file refdna_file information to be used to calculate RNA VAF (Default=NA).

See "https://github.com/hase62/Neoantimon"

samtools_dir The file directory to samtools_0_x_x (Default="samtools"). It shouled be indicated when you indicate RNA-bam and try to calculate RNA VAF.

bcftools_dir The file directory to netMHCpan (Default="bcftools"). It shouled be indicated when you indicate RNA-bam and try to calculate RNA VAF . samtools 0_x_x includes bcftools in the directory.

chr_column The column number describing chromosome number in input_file (Default=NA, but will automatically search "Chr" in header).

mutation_start_column

cnv_file

The column number describing mutation start Position in input_file (Default=NA, but will automatically search "Start" in header).

mutation_end_column

The column number describing mutation end Position in input_file (Default=NA, but will automatically search "End" in header).

mutation_ref_column

The column number describing mutation Ref in input_file (Default=NA, but will automatically search "Ref" in header).

 $\verb|mutation_a| t_b \verb|nd_column|$

The column number describing mutation Alt (BND format) in input_file (Default=NA, but will automatically search "Alt" in header).

depth_normal_column

 $\label{thm:column} The \ column \ number \ describing \ the \ read \ count \ from \ normal \ cells \ (Default=NA).$ $\ depth_tumor_column$

The column number describing the read count from tumor cells (Default = NA).

nm_id_column (Required if gene_symbol_column = NA) The column number describing NM IDs in input_file such as

"SLCO1C1:NM_001145944:exon7:c.692_693insG:p.L231fs" (Default=NA).

ambiguous_between_exon

The maximum number to permit the differences between Exon-Lengths from refFlat and refMrna (Default=0).

ambiguous_codon

The maximum number to permit the differences between inputfile- and refMrnaoriented translation start/end position (Default=0). MainSVFUSIONClass2 37

peptide_length Peptide Length to be generated (Default = 15 in HLA Class2).
gene_symbol_column

(Required if nm_id_column = NA) The column number describing gene symbol

in input_file (Default=NA).

mate_id_column (Required) The column indicating mateIDs or svIDs such as "SVMERGE1_1"

(Default=NA).

ignore_short Ignore to output results of short peptide less than min (peptide_length)

Value

void (Calculated Neoantigen Files will be generated as .tsv files.):

HLA: HLA type used to calculate neoantigen.

Pos: The position of a fraction of peptide used to be evaluated from the full-length peptide.

Gene Gene symbol used to be evaluated in NetMHCpan.

Evaluated_Mutant_Peptide: The mutant peptide to be evaluated.

Evaluated_Mutant_Peptide_Core: The core peptide of the mutant peptide to be evaluated in NetMHC-pan.

Mut IC50: IC50 value for evaluated mutant peptide.

Mut_Rank: Rank value for evaluated mutanat peptide.

Chr: Chromosome Number of the mutation.

NM_ID: NM_ID used to construct peptides from the mutation.

Change: The annotation to be described in .vcf file.

Ref: reference type nucleic acid base.

Alt: alternative type nucleic acid base.

Prob: A probability of reference nucleic acid base described in .vcf file.

Mutation_Prob: A probability of alternative nucleic acid base described in .vcf file.

Exon_Start: The exon start position of the corrsponding NM_ID.

Exon_End: The exon end position of the corrsponding NM_ID.

Mutation_Position: The mutation position of the corrsponding NM_ID.

Total_Depth: The sum depth of the reference and alternative nucleic acid base.

Tumor_Depth: The depth of the alternative nucleic acid base.

Wt_Peptide: The full-length of the wild-type peptide.

Mutant_Peptide: The full-length of the mutant peptide.

Total_RNA: The expression amount of the corresponding RNA.

Tumor_RNA_Ratio: The variant allele frequency of the corresponding RNA.

Tumor_RNA: The modified amount of the corresponding RNA level based on RNA Reads.

Tumor_RNA_based_on_DNA: The modified amount of the corresponding RNA level based on DNA Reads.

MutRatio: The mean value of the cancer cell fraction probability.

MutRatio_Min: The 1% percentile of the cancer cell fraction probability.

MutRatio_Max: The 99% percentile of the cancer cell fraction probability.

P_I: Priority score using the IC50. Please use CalculatePriorityScores <- function(result, useR-NAvaf = FALSE)

38 sample_hla_table_c2

P_R: Priority score using the percentage of rank affinity. Please use CalculatePriorityScores <-function(result, useRNAvaf = FALSE)

P: Priority score implemented in MuPeXI (Bjerregaard et al. 2017). Please use CalculatePriorityScores <- function(result, useRNAvaf = FALSE)

sample_copynum

A Format / Sample file for Copy Number Information

Description

A dataset containing the copy number information obtained by, e.g., ASCAT.

Usage

```
data(sample_copynum)
```

Format

A data frame with 7 rows and 9 variables

Description

A dataset containing the HLA types of patients in each row.

Usage

```
data(sample_hla_table_c1)
```

Format

A data frame with 3 rows and at most 7 variables

Description

A dataset containing the HLA types of patients in each row.

Usage

```
data(sample_hla_table_c2)
```

Format

A data frame with at least 3 row and at most 10 variables

sample_refFlat.grch37 39

```
{\tt sample\_refFlat.grch37} \ \ \textit{A Sample file for refFlat}
```

Description

A dataset containing a part of refFlat data.

Usage

```
data(sample_refFlat.grch37)
```

Format

A data frame with 11 column.

```
sample_refMrna.grch37.fa A \ \textit{Sample file for refSeq RNA}
```

Description

A dataset containing a part of refSeq RNA.

Usage

```
data(sample_refMrna.grch37.fa)
```

Format

A data frame with 1 column.

```
sample\_result\_INDEL\_CLASS1\_ALL\\ Analyzed\ Result\ for\ INDEL\ CLASS1
```

Description

Analyzed Result for INDEL CLASS1

Usage

```
data(sample_result_INDEL_CLASS1_ALL)
```

```
sample_result_INDEL_CLASS2_ALL

Analyzed Result for INDEL CLASS2
```

Description

Analyzed Result for INDEL CLASS2

Usage

```
{\tt data(sample\_result\_INDEL\_CLASS2\_ALL)}
```

```
sample_result_SeqFragment_CLASS1_ALL

Analyzed Result for A DNA Fragment CLASS1
```

Description

Analyzed Result for A DNA Fragment CLASS1

Usage

```
data(sample_result_SeqFragment_CLASS1_ALL)
```

```
sample_result_SeqFragment_CLASS2_ALL

Analyzed Result for A DNA Fragment CLASS2
```

Description

Analyzed Result for A DNA Fragment CLASS2

Usage

```
data(sample_result_SeqFragment_CLASS2_ALL)
```

```
sample\_result\_SNV\_CLASS1\_ALL\\ Analyzed\ Result\ for\ SNV\ CLASS1
```

Description

Analyzed Result for SNV CLASS1

Usage

```
data(sample_result_SNV_CLASS1_ALL)
```

```
sample_result_SNV_CLASS2_ALL
```

Analyzed Result for SNV CLASS2

Description

Analyzed Result for SNV CLASS2

Usage

```
data(sample_result_SNV_CLASS2_ALL)
```

```
sample_result_SVFusion_CLASS1_ALL
```

Analyzed Result for SV Fusion CLASS1

Description

Analyzed Result for SV Fusion CLASS1

Usage

```
data(sample_result_SVFusion_CLASS1_ALL)
```

```
sample_result_SVFusion_CLASS2_ALL
```

Analyzed Result for SVFusion CLASS2

Description

Analyzed Result for SVFusion CLASS2

Usage

```
data(sample_result_SVFusion_CLASS2_ALL)
```

sample_rna_exp

A Format / Sample file for RNA Expression Information

Description

A dataset containing the RNA expression amount of patient for each gene.

Usage

```
data(sample_rna_exp)
```

Format

A data frame with 22 rows and 3 variables

42 sample_vcf.snps

sample_sv_bnd

A Format / Sample file for Annotated vcf file.

Description

A dataset containing the variant information of a patient.

Usage

```
data(sample_sv_bnd)
```

Format

A data frame with 9 rows and variables including "Chr" "Start" "End" "Ref" "Alt (BND format)" "Func.refGene (exonic, intron, intergenic, ...)" "ExonicFunc.refGene (exonic nonsynonymous, synonymous, insertion, ...)" "mateID (e.g., SVMERGE1_1)"

sample_vcf.annovar

A Format / Sample file for Annotated vcf file basef on Annovar.

Description

A dataset containing the variant information of a patient.

Usage

```
data(sample_vcf.annovar)
```

Format

A data frame with 9 rows and variables including "Chr" "Start" "End" "Ref" "Alt" "Func.refGene (exonic, intron, intergenic, ...)" "ExonicFunc.refGene (exonic nonsynonymous, synonymous, insertion, ...)" "AAChange.refGene (e.g., SLCO1C1:NM_001145944:exon7:c.692_693insG:p.L231fs ...)"

sample_vcf.snps

A Format / Sample file for snp informatin.

Description

A dataset containing snps information of a patient.

Usage

```
data(sample_vcf.snps)
```

Format

A data frame with variables including "#CHROM" "POS" "ID" "REF" "ALT" "QUAL" "FILTER" "INFO" "FORMAT".

sample_vcf.vep 43

sample_vcf.vep

A Format / Sample file for Annotated vcf file based on VEP.

Description

A dataset containing the variant information of a patient.

Usage

```
data(sample_vcf.vep)
```

Format

A data frame with variables including "#Uploaded_variation" "Location" "Allele" "Gene" "Feature" "Feature_type" "Consequence" "cDNA_position" "CDS_position" "Protein_position" "Amino_acids Codons" "Existing_variation" "Extra"

TestAnalysis

Execute Sample Analysis

Description

Execute Sample Analysis

Usage

TestAnalysis()

Value

void

Index

```
*Topic datasets
                                                sample_hla_table_c1, 38
    sample_copynum, 38
                                                sample_hla_table_c2, 38
    sample_hla_table_c1, 38
                                                sample_refFlat.grch37, 39
    sample_hla_table_c2, 38
                                                sample_refMrna.grch37.fa, 39
    sample_refFlat.grch37,39
                                                sample_result_INDEL_CLASS1_ALL, 39
    sample_refMrna.grch37.fa, 39
                                                sample\_result\_INDEL\_CLASS2\_ALL, 40
    sample_rna_exp, 41
                                                sample_result_SeqFragment_CLASS1_ALL,
    sample_sv_bnd, 42
    sample_vcf.annovar, 42
                                                sample_result_SeqFragment_CLASS2_ALL,
    sample_vcf.snps, 42
    sample_vcf.vep, 43
                                                sample_result_SNV_CLASS1_ALL, 40
*Topic result
                                                sample_result_SNV_CLASS2_ALL, 41
    sample_result_INDEL_CLASS1_ALL, 39
                                                sample_result_SVFusion_CLASS1_ALL, 41
    sample_result_INDEL_CLASS2_ALL, 40
                                                sample_result_SVFusion_CLASS2_ALL, 41
    sample_result_SeqFragment_CLASS1_ALL,
                                                sample_rna_exp, 41
                                                sample_sv_bnd, 42
    sample_result_SeqFragment_CLASS2_ALL,
                                                sample_vcf.annovar, 42
                                                sample_vcf.snps, 42
    sample_result_SNV_CLASS1_ALL, 40
                                                sample_vcf.vep, 43
    sample_result_SNV_CLASS2_ALL, 41
                                                TestAnalysis, 43
    {\tt sample\_result\_SVFusion\_CLASS1\_ALL},
    sample_result_SVFusion_CLASS2_ALL,
        41
CalculatePriorityScores, 2
Export_Summary_Entire_Fragments, 3
Export_Summary_Fragments, 4
Export_Summary_IndelSV, 5
Export_Summary_IndelSV_perFragments, 6
Export_Summary_SNV, 7
MainEntireRegionClass1,8
MainEntireRegionClass2, 10
MainINDELClass1, 12
MainINDELClass2, 15
MainSeqFragmentClass1, 19
MainSeqFragmentClass2, 21
MainSNVClass1, 23
MainSNVClass2, 27
MainSVFUSIONClass1, 31
MainSVFUSIONClass2, 34
sample_copynum, 38
```