# Package 'IrRNAseqBenchmark'

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```
Type Package
Title Figure out how to call varaints from Iso-Seq data
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 ${\bf URL} \ {\tt https://github.com/vladimirsouza/lrRNAseqBenchmark}$ 

 $\pmb{BugReports} \ \text{https://github.com/vladimirsouza/lrRNAseqBenchmark/issues}$ 

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 $add\_a\_method\_indel\_information\_to\_master\_table$ 

Add column that states whether the variant called by a method is an indel or not

## Description

Add column that states whether the variant called by a method is an indel or not

## Usage

```
add_a_method_indel_information_to_master_table(
  input_table,
  vcf_file,
 method_name,
  truth_name
)
```

## **Arguments**

input\_table A data.frame. The master table to add the new column. vcf\_file 1-length string. The address of the method VCF file.

method\_name 1-length string. The name of the method.

truth\_name 1-length string. The name of the ground-truth.

#### Value

A data.frame

```
add_format_tag_from_vcf
                         Add a FORMAT tag from a VCF file to a master table
```

## **Description**

The function adds to 'input\_table' the new column <tagName>\_<methodName>, where <tag-Name> is 'tag' in lower case and <methodName> is 'method\_name'.

## Usage

```
add_format_tag_from_vcf(input_table, method_name, tag, vcf_file)
```

## **Arguments**

input\_table A data.frame. The input master table.

method\_name A 1-length string. The name of the method. tag

A 1-length string. The name of the tag to add. It must be exactly how the tag is

written in the VCF file (case sensitive).

vcf\_file A 1-length string. The path to the VCF file from which the tag is extracted.

#### Value

A data.frame.

```
add_homopolymer_length_when_indels

Add homopolymer lengths to a master table
```

## **Description**

This function adds to the master table a column with the lengths of homopolymers, from the reference fasta, that overlaps positions POS+1, where POS is the position of a variant. POS+1 makes sence because minimp2 places homopolymer indels to the left of homopolymers, and , in case of indels, the column POS of VCF files means the position immediately to the left of the indel. In this way, homopolymer lengths of SNPs are meaningless. Moreover, homopolymer lengths of variants that are heterozygous alternatives should be meaningless as well. That is because they could contain alleles that are SNPs.

## Usage

```
add_homopolymer_length_when_indels(
  input_table,
  homopolymers,
  ouput_what = "length"
)
```

## **Arguments**

input\_table A data.frame. The master table to add the new column.

homopolymers A CompressedIRangesList object. It should store all homopolymers, it's nucleo-

tive types and lengths, of the genome used as the reference to call the variants.

It is gerated by the function 'sarlacc::homopolymers'.

ouput\_what A string equal to "length" (default) or "nts". If "length", the lengths of ho-

mopolymers are ouput (1 for non-homopolymers). If "nts", the nucleotide type

is output (NA for non-homopolymers).

## Value

A data.frame

## **Description**

If a variant in the VCF file doesn'r contain the specified tag, the function return -1 for that variant.

```
add_info_tag_from_vcf(input_table, col_name = NULL, tag, vcf_file)
```

input\_table A data.frame. The input master table.

col\_name A 1-length string. the name of the column to add. If NULL (default), 'col\_name'

is equal to 'tag'.

tag A 1-length string. The name of the tag to add. It must be exactly how the tag is

written in the VCF file (case sensitive).

vcf\_file A 1-length string. The path to the VCF file from which the tag is extracted.

## Value

A data.frame.

```
add_method_vs_truth_comparison_to_master_table
```

Add classiffication of variants calls by comparing to the ground-truth to a master table

## **Description**

Compare variants calls of a method to the ground-truth and classify each one as true-positiove (TP), false-negative (FN), false-positive (FP), or true-negative (TN).

## Usage

```
add_method_vs_truth_comparison_to_master_table(
  input_table,
  method_name,
  truth_name,
  replace_column = FALSE
)
```

## **Arguments**

input\_table A data.frame. The master table to add the new column.

method\_name A 1-length string. The name of the method to compare to the ground-truth.

truth\_name A 1-length string. The name of the ground-truth.

replace\_column 1-length boolean (default is FALSE). If the classification column already exists

and it is desired to replace it.

## Value

A data.frame.

6 add\_qual\_from\_vcf

```
add_number_of_n_cigar_reads_to_master_table

Add the count of N-cigar reads to the master table
```

## **Description**

This function adds a new column to a master table with the counts of N-cigar reads that overlap each site.

## Usage

```
add_number_of_n_cigar_reads_to_master_table(
  input_table,
  input_bam,
  dataset_name = NULL
)
```

## Arguments

input\_table A data.frame. The master table to add the new column.

input\_bam A 'GenomicAlignments' object from which N-cigar-read counts are got.

dataset\_name A 1-length string used to set the name of the new column.

#### Value

A data.frame.

add\_qual\_from\_vcf

Add QUAL from a VCF into a master table

## **Description**

Add QUAL to master table.

## Usage

```
add_qual_from_vcf(input_table, method_name, vcf_file)
```

## **Arguments**

input\_table A data.frame. The master table to add the new column.

method\_name A 1-lenght string. The name of the method from which is desirable to get the

QUAL values. The new column is named as "qual\_<method\_name>".

vcf\_file A 1-lenght string. The path of the VCF file from which the the QUAL values

are extracted.

## Value

A data.frame

```
add_read_coverage_from_bam_to_master_table

Add read coverage (taken from BAM file) of each dataset to a master table
```

## **Description**

Add read coverage (taken from BAM file) of each dataset to a master table

## Usage

```
add_read_coverage_from_bam_to_master_table(
  input_table,
  dataset_coverage,
  dataset_name
)
```

## **Arguments**

```
input_table A master table to add read-coverage columns.

dataset_coverage

Object generated by IRanges::coverage function.

dataset_name A 1-length string. Name of the dataset used to genere 'dataset_coverage'.
```

## Value

A data frame.

```
add_some_read_statistic_from_bam_metacolumn

Add some statistic of a tag about reads of a BAM file
```

## **Description**

Add some statistic of a tag about reads of a BAM file

```
add_some_read_statistic_from_bam_metacolumn(
  input_table,
  col_name,
  galn,
  tag,
  stat
)
```

input\_table A data.frame. The input master table.

col\_name A 1-length string. The name of the new column to add.

galn A GAlignment object. The alingnment from where the tag is extracted.

tag A 1-length string. The name of tag that is wanted to calculate the statistic 'stat'.

The name must be exactly the same of one column name from 'mcols(galn)'.

stat A function used to calculate the desired statistic.

#### Value

A data.frame.

```
add_splice_site_info_to_master_table
```

Add columns about spice sites to a master table

## **Description**

Add columns about spice sites to a master table

## Usage

```
add_splice_site_info_to_master_table(
  input_table,
  splice_sites,
  max_dist_from_splice_site = 20,
  multiply_max_dist = 100
)
```

## **Arguments**

A 1-length integer (default is 20) that indicates the maximum distance of a variant from any splice site to consider it a variant near a splice site.

multiply\_max\_dist

A 1-length integer (default is 100). To create the column 'ss\_shortest\_dist' from column 'ss\_dist', NA values (which means variables far from any splice site) are converted to integers. To make these integers be a high value, they are set to max\_dist\_from\_splice\_site \* multiply\_max\_dist.

## Value

A data.frame.

```
{\it add\_the\_ground\_truth\_indel\_information\_to\_master\_table} \\ {\it Add\ column\ that\ states\ whether\ the\ variant\ in\ the\ ground\_truth\ is\ an\ indel\ or\ not}}
```

## **Description**

Add column that states whether the variant in the ground-truth is an indel or not

## Usage

```
add_the_ground_truth_indel_information_to_master_table(
  input_table,
  vcf_file,
  truth_name
)
```

## **Arguments**

```
input_table A data.frame. The master table to add the new column.

vcf_file 1-length string. The address of the ground-truth VCF file.

truth_name 1-length string. The name of the ground-truth.
```

## Value

A data.frame

```
add_two_method_comparison_to_master_table

Add column to compare two methods in a master table
```

## Description

This function adds a column in a master table that compares two given methods. The new column informs whether the variant was called only by the first method ('method1\_name'), only by the second method ('method2\_name'), by both methods ('"both"'), or neither of them ('"neither"').

```
add_two_method_comparison_to_master_table(
  input_table,
  method1_name,
  method2_name
)
```

input\_table A data.frame. The master table to add the new column.

method1\_name A 1-length string. The name of the first method.

method2\_name A 1-length string. The name of the second method.

#### Value

A data.frame.

 $add\_variant\_density\_of\_a\_method$ 

Add column of density of variants around each variant called by a method

## Description

Add column of density of variants around each variant called by a method

## Usage

```
add_variant_density_of_a_method(input_table, window_size, used_methods)
```

## Arguments

input\_table A data.frame. The input master table.

window\_size A 1-length integer. The size of the window where variants are used to calculate

the density of variants around a each variant called by the specified method.

 $\mbox{used\_methods} \qquad \mbox{A vector of strings. The names of the methods to be used.}$ 

## Value

A data.frame.

```
add_variant_type_to_master_table
```

Add a column that states the variant type according a method or the ground truth

## **Description**

The new column stores the values "snp", "insertion", or "deletion", respectively to the variant type. If the variant type can not be defined – because the variant is heterozygous alternative and the alleles are a mix between SNP, insertion, or deletion – the value returned is "mix".

## Usage

```
add_variant_type_to_master_table(input_table, vcf_file, method_name)
```

## **Arguments**

```
input_table A data.frame. The master table to add the new column.vcf_file 1-length string. The address of the VCF file.method_name 1-length string. The name of the ground-truth or method.
```

#### Value

A data.frame

```
calculate_precision_recall_for_multi_master_tables

Calculate accuracy measures for multiple master tables
```

## Description

Create a data.frame with accuracy measures for multiple master tables.

```
calculate_precision_recall_for_multi_master_tables(
    ...,
    experiment_names,
    method_names,
    output_method_names = NULL,
    data_names,
    truth_names,
    coverage_thresholds = c(3, 15, 40, 100),
    what
)
```

.. Master tables.

experiment\_names

Vector of strings. Name of the experiments in the same order of the master

tables.

method\_names

Vector or list of strings. Names of the methods to compare. If all experiments compare the same methods, 'method\_names' can be a single vector. Otherwise, 'method\_names' must be a list, in which each element specifies the names of the methods to compare for each experiment.

output\_method\_names

NULL (default), or a vector or a list of strings. Names of the methods to output.

data\_names Vector or list of strings.
truth\_names Vector or list of strings.

coverage\_thresholds

Vector or list of integers.

what Vector or list of strings.

#### Value

A data.frame.

```
calculate_precision_recall_for_n_cigar_read_count_intervarls
```

For each N-cigar-read-count interval, calculate accuracy measures for multiple master tables

## **Description**

Create a data.frame with accuracy measures for multiple master tables and different intervals for N-cigar-read counts.

```
calculate_precision_recall_for_n_cigar_read_count_intervarls(
    ...,
    experiment_names,
    method_names,
    output_method_names = NULL,
    data_names,
    truth_names,
    start_n_cigar_read_percent_intervals = c(0, 0.05, 0.9),
    end_n_cigar_read_percent_intervals = c(0.05, 0.9, 1),
    what
)
```

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#### **Arguments**

. Master tables.

experiment\_names

Vector of strings. Name of the experiments in the same order of the master

tables.

method\_names Vector or list of strings. Names of the methods to compare. If all experiments

compare the same methods, 'method\_names' can be a single vector. Otherwise, 'method\_names' must be a list, in which each element specifies the names of

the methods to compare for each experiment.

output\_method\_names

NULL (default), or a vector or a list of strings. Names of the methods to output.

data\_names Vector or list of strings.
truth\_names Vector or list of strings.
start\_n\_cigar\_read\_percent\_intervals

vector of doubles. The start of each interval of N-cigar-read counts.

end\_n\_cigar\_read\_percent\_intervals

vector of doubles. The end of each interval of N-cigar-read counts.

what Vector or list of strings.

#### Value

A data.frame.

calc\_accuracy\_measures

Calculate precision, sensitivity and F1-score of a method in a master table

## **Description**

Calculate precision, sensitivity and F1-score of a method in a master table

## Usage

```
calc_accuracy_measures(input_table, method_name, truth_name)
```

## **Arguments**

input\_table A data.frame. The master table.

method\_name A 1-length string. The name of the method to calculate the accuracy measures.

truth\_name A 1-length string. The name of the ground-truth to validate the method.

#### Value

A named vector with the accuracy measures.

```
check_accuracy_per_coverage
```

Barplot of accuracy measures for different methods and coverage

## **Description**

Return a ggplot object for visualization.

## Usage

```
check_accuracy_per_coverage(
  master_table,
  method_names,
  output_method_names = NULL,
  data_name,
  truth_name,
  coverage_threshold
)
```

## **Arguments**

master\_table A data.frame. The input master table.

method\_names A vector of strings. The names of the methods to be compared in the master

table.

output\_method\_names

A vector of strings. How to output the names of the methods to be compared.

The defaut is NULL.

data\_name A 1-length string. The name of the dataset used with the methods to be com-

pared.

truth\_name A 1-length string. The name of the ground-truth.

coverage\_threshold

A vector of integers. The minimum thresholds to filer by read coverage.

## Value

A ggplot object.

get\_splice\_sites\_info

get\_splice\_sites\_info Get all splice sites positions of a BAM files

## Description

Get the position of all splice sites in a BAM file and the number of reads that support each one of them. Besides that, indicate whether the splice site is acceptor or donor site.

## Usage

```
get_splice_sites_info(input_bam, threads)
```

## Arguments

input\_bam The input BAM file to extract splice site positions from.

threads Number of threads.

## **Details**

This function may take several hours to run.

## Value

A data.frame in which each row is a splice-site position.

gt_vt_method	get the genotype and the variant type of the calls of a primary and secundary method
--------------	--

## Description

Create a new data.frame from a input master table. The rows correspond to the same variants in 'input\_table' and in the same order.

```
gt_vt_method(input_table, gt_first, gt_second, vt_first, vt_second)
```

input\_table A data.frame. The input master table.

gt\_first A 1-length string. The name of the column that stores the genotypes called by a method. This method is called the first method.

gt\_second A 1-length string. If the first method does not call some variants, extract their genotype from column 'gt\_second'.

vt\_first A 1-length string. The name of the column that stores the variant type called by a method. This method is called the first method.

vt\_second A 1-length string. If the first method does not call some variants, extract their

variant type from column 'vt\_second'.

#### Value

A data.frame.

igv\_batch\_screenshots Generate igv batch screenshots script

## **Description**

Generate igv batch screenshots script

## Usage

```
igv_batch_screenshots(
  chrm,
  pos,
  output_dir,
  prefix,
  snapshot_path,
  windows_size = 1501,
  screenshot_number = 100,
  output_positions = FALSE
)
```

## **Arguments**

chrm A vector of string. Chromossome name, e.g. "chrm1".

pos A vector of integers. Positions to center the screenshot.

output\_dir A 1-length string. Directory where the IGV-batch-screenshot script is saved.

prefix A 1-length string.

snapshot\_path A 1-length string. Directory where the IGV-batch-screenshot script will save the

screenshots.

windows\_size 1-length integer. Window size of the screenshot.

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```
screenshot_number
```

1-length integer. Number of screenshot to generate.

output\_positions

If TRUE, output a data.frame with variant positions and the reagions to visualize.

initiate\_master\_table Initiate master table

## **Description**

This function create the first columns of the master table. Those columns indicate whether a method could call of not a variant. A master table is used to get information form VCF files generated from different methods and compare to ground-truth VCF files.

## Usage

```
initiate_master_table(..., method_names)
```

## **Arguments**

... VCF file addresses.

method\_names

Vector of strings in which each element is the identification (name) of each input VCF files. The order of these elements must be in accordance with the order of the input files, and its length must be the same as the number of input VCF files.

## Value

A data.frame.

is\_indel\_method

Add a column to state whether the variant is an indel or not

## **Description**

To classify the variant as an indel, first the function looks at the VCF file of the 'first\_method'. If the variant is not there, if looks at the VCF file of the 'second\_method'.

#### Usage

```
is_indel_method(input_table, first_method, second_method = NULL)
```

## **Arguments**

input\_table A data.frame. The master table to add the new column.

first\_method A 1-length string. The name of the first method. second\_method A 1-length string. The name of the second method

#### **Details**

Output meaning: \* '-1' means that the variant type couldn't be defined, because it is heterozygous alternative; \* '0' means it is not an indel; \* '1' means it is an indles; \* 'NA' means the 'first\_method' (and the 'second method') didn't call the variant.

#### Value

A data.frame.

make\_homopolymer\_plot Make plot to compare accuracy per homopolymer length

## **Description**

The function uses the output from function 'method\_homopolymer\_indels' to make a plot to compare how the accuracy vary according to the homopolymer length or not within homopolymers. Those accuracy measures can be either rates of TPs, FNs and FPs, or the precision, recall and F1-score.

## Usage

```
make_homopolymer_plot(
  input_hom_table,
  variant_type,
  method_name,
  truth_name,
  hom_length_intervals,
  interval_names,
  to_calculate
)
```

## **Arguments**

input\_hom\_table

A data.frame. The output of the function 'method\_homopolymer\_indels'.

variant\_type

A 1-length string. The variant type. Possivle values are: "snp", "deletion", or

"insertion".

method\_name

A 1-length string. The name of the method to analyse.

truth\_name

A 1-length string. The name of the ground-truth.

hom\_length\_intervals

A vector of integers. Must be the same length of 'interval\_names'. The inferior

limit for each interval for homopolymer length.

interval\_names A vector of strings. Must be the same length of 'hom\_length\_intervals'. The

name of each interval of homopolymer length.

to\_calculate

A 1-length string. Possible values are: "rates" or "pre\_rec\_f1".

#### Value

A list, in which the first element, named 'p', is a ggplot object, and the second element, named 'interval\_counts', is a named vector of integers that stores the counts of variants in each interval specified by the x-axis of the plot in 'p'.

```
make_homopolymer_table_to_plot
```

Organize the data used to make plots about homopolymer analysis

## **Description**

The user may want to use this function several times to pull information about different combinations between 'variant\_type' and 'method\_name' from 'input\_hom\_table'. In the future, the function should make the job automatically using loops.

## Usage

```
make_homopolymer_table_to_plot(
   input_hom_table,
   variant_type,
   method_name,
   truth_name,
   hom_length_intervals,
   interval_names,
   to_calculate,
   output_method_name
)
```

## **Arguments**

input\_hom\_table

A data.frame generated by function 'method\_homopolymer\_indels'.

variant\_type A 1-length string. Possible values are "insertion" or "deletion".

method\_name A 1-length string. The name of the method from which is desired to extract

information.

truth\_name A 1-length string. The name of the ground-truth.

hom\_length\_intervals

A vector of integers. The minimum for each interval of homopolymer length. Each interval 'i' ranges from 'hom\_length\_intervals[i]' to 'hom\_length\_intervals[i+1]', except the last interval which upper limit is 'Inf'.

interval\_names A vector of characters with the same length of 'hom length intervals'. The

name for each interval of homopolymer length.

to\_calculate A 1-length string. Possible values are "rates" or "pre\_rec\_f1". If "rates", the

functin calculates the rates of TPs, FNs and FPs. If "pre\_rec\_f1", it calculates

the precision, the recall and the F1-score.

```
output_method_name
```

A 1-length string. The label of the method specified in 'method\_name' to be output.

## Value

```
A 2-length list ('class_counts' and 'dat_text').
```

```
method_homopolymer_indels
```

Get indels in homopolymers

## **Description**

Create a new data.frame that stores deletions of inside homopolymers, and insertions of a same nucleotide type that happen inside homopolymers of that same nucleotide type. Homopolymer length equal to 1 means non-homopolymers.

## Usage

```
method_homopolymer_indels(
   input_table,
   first_method_name,
   second_method_name,
   vcf_first,
   vcf_second,
   homopolymers,
   ref_fasta_seqs,
   min_isoseq_coverage,
   genotyped_alt
)
```

## **Arguments**

```
input_table A data.frame. The input master table.
```

first\_method\_name

A 1-length string of the name of a method. First, the variant information that is taken is related to this method. If a variant is not called by the method, its information is taken relating to the method specifiec in 'second\_method\_name'.

second\_method\_name

A 1-length string. The name of a method.

vcf\_first A 1-length string. The path of the VCF file of the first method.

vcf\_second A 1-length string. The path of the VCF file of the second method.

homopolymers A CompressedIRangesList object. It should store all homopolymers, it's nucleotive types and lengths, of the genome used as the reference to call the variants.

It is gerated by the function 'sarlacc::homopolymerFinder'.

ref\_fasta\_seqs A DNAStringSet object. The sequences of the genome used as the reference to call the variants. It's names must be in the form like "chr1", "chr2", ..., "chrX", "chrY".

min\_isoseq\_coverage

min iso-seq read coverage to filter.

genotyped\_alt

One of the strings "find" or "same". If "find", the function finds the correct alternative allele based on the genotype. This option must be chosen if there are multiple values for column ALT, which is the case of VCF files output by Deep-Variant and Clair3. Since GATK's VCF files keep only the genotyped alternative allele in column ALT, this argument should be "same".

#### Value

A data.frame.

```
precision_recall_curve_per_coverage
```

Precision-recall curves to compare methods, using different filtering on read coverage

## **Description**

Precision-recall curves to compare methods, using different filtering on read coverage

## Usage

```
precision_recall_curve_per_coverage(
  master_table,
  method_names,
  output_method_names = NULL,
  data_name,
  truth_name,
  coverage_thresholds,
  what
)
```

## Arguments

master\_table A data.frame. The input master table.

method\_names A vector of strings. The names of the methods to be compared.

output\_method\_names

A vector of strings. How to output the names of the methods to be compared.

The default is NULL.

data\_name A 1-length string. The name of the dataset used with the methods to be com-

pared.

truth\_name A 1-length string. The name of the ground-truth.

coverage\_thresholds

A vector of integers. The minimum thresholds to filer by read coverage. Each

element defines a point in the precision-recall curves.

A 1-length string. Possible values are: "snps\_indels" (default), to make curves

for SNPs and indels separately; "snps", to make curves only for SNPs; "indels", to make curves only for indels; "overall", to make curves without distinguishing

SNPs and indels.

## Value

what

A ggplot object.

```
splice_junction_analysis_table
```

Generate data for splice-junction analysis using multiple master tables

## **Description**

Generate data to plot variant performance of sites near to and far from splice junctions comparisons using multiple master table as facets.

## Usage

```
splice_junction_analysis_table(
    ...,
    experiment_names,
    truth_names,
    method_names,
    output_method_names,
    variant_type,
    min_isoseq_coverage
)
```

## **Arguments**

... 1-length strings. Paths to master tables saved as RDS files.

experiment\_names

A vector of strings with length equal to the number of master tables input in '...'.

Names of the datasets for each data table.

truth\_names A vector of strings. Names of the ground truth in each data table especified in

·...'.

method\_names A vector of strings with length equal to the number of master tables input in '...'.

Names of the methods to compare.

output\_method\_names

A vector of strings with length equal to the number of master tables input in '...'. Names of the methods to be in the output.

standardize\_genotype 23

```
\label{lem:condition} \begin{tabular}{ll} variant\_type & A 1-length string. Possible values are "snp" or "indel". \\ min\_isoseq\_coverage \\ \end{tabular}
```

A 1-length integer. The threshod value for the minimum Iso-Seq read coverage.

#### Value

A list of two elements ('acc\_sj' and 'n\_test') to be used to draw a chart to analyse the variant-calling performance of sites near to and far from splice junctions. 'acc\_sj' stores the calculated performance measures, and 'n\_test' stores the number of observed variants in each case.

standardize\_genotype Standardize genotypes

## Description

```
This function rewrite genotype like: * 0|1 to 0/1; * 0/2, 0/3, ..., to 0/1; * 2/2, 3/3, ..., to 1/1; * 1/3, 2/3, ..., to 1/2
```

## Usage

```
standardize_genotype(gt)
```

## Arguments

gt

A vector of strings. The input genotypes.

#### Value

A vector of strings.

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