Package 'SaTAnn'

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|--|
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| annotate_ORFs Annotate detected ORFs in transcript and genome space | | | | | | | | | |

Description

This function annotates quantified ORFs with respect to other detected ORFs and annotated ones, in both genome and transcript space.

Usage

```
annotate_ORFs(results_ORFs, Annotation, genome_sequence, region,
  genetic_code)
```

Arguments

results_ORFs Full list of detected ORFs, from select_quantify_ORFs

Annotation Rannot object containing annotation of CDS and transcript structures (see prepare_annotation_files genome_sequence

BSgenome object

region genomic region being analyzed genetic_code GENETIC_CODE table to use

Details

As multiple transcripts can contain the same ORF, all the transcript and transcript biotypes are indicated, with a preference for protein_coding transcripts in the "compatible" columns (to be conservative when assessing translation of non-protein coding transcripts). Such compatibility is also output considering the most upstream start codon for that ORF.

Splice features of each orf is annotated with respect to the longest coding transcripts and to the highest translated ORF in that gene.

Variants in N or C terminus of the translated proteins are also indicated (Beta).

ORF annotation with respect to the annotated transcript is also indicated, as follows:

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novel: no ORF annotated in the transcript.

ORF_annotated: same exact ORF as annotated.

N_extension: N terminal extension. N_truncation: N terminal extension.

uORF: upstream ORF.

overl_uORF: upstream overlappin uORF. NC_extension: N and C termini extension.

dORF: downstream ORF.

overl_dORF: downstream overlapping ORF.

nested_ORF: nested ORF.

C_truncation: C terminal truncation. C_extension: C terminal extension.

As transcipt-specific annotation can be misleading due to a plethora of different transcripts, it is important to distinguish ORFs also on the basis of their overlap with know CDS regions. ORF annotation with respect to the entire set of CDS exon for the analyzed genomic regions is indicated as follows:

novel: No CDS region is annotated in the entire region.

novel_Upstream: ORF is upstream of annotated CDS regions (does not overlap).

novel_Downstream: ORF is downstream of annotated CDS regions (does not overlap).

novel_Internal: genomic location of the ORF is present between the start of the first, and the end of the last CDS region (does not overlap).

exact_start_stop: Same start and end locations.

Alt5_start: Different start region, upstream.

Alt3_start: Different start region, downstream.

Alt5_stop: Different end region, upstream.

Alt3_stop: Different end region, downstream.

Another layer of annotation is performed by checking the position of the ORF stop codon with respect to the last exon-exon junction.

Value

Exon structure of detected ORF including possible missing exons from reference, together with a spl_type column including the annotation for each exon (e.g. alternative acceptors or donor).

Additional columns are added to the ORFs tx object:

compatible_with: Set of transcript ids possibly containing the entire ORF structure.

compatible_biotype: Compatible transcript biotype; if a protein coding transcript can contain the ORF, this is set to protein_coding.

compatible_tx: One selected compatible transcript (preference if protein_coding).

compatible_ORF_id_tr: ORF_id_tr id if selecting the compatible transcript.

compatible_with_longest: Same as compatible_with but using the most upstream start codon. compatible_ORF_id_tr_longest: Same as compatible_ORF_id_tr but using the most upstream start codon .

ref_id: transcript_id of the transcript used to annotate splicing (longest).

ref_id_maxORF: ORF_id_tr of the ORF used to annotated splicing (most translated of the gene).

NC_protein_isoform: Annotation of possible N or C termini variant (when transcript is pro-

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```
tein_coding).
```

ORF_category_Tx: ORF annotation with respect to ORF position in the transcript .

ORF_category_Tx_compatible: ORF annotation with respect to ORF position in the transcript, using the compatible_ORF_id_tr.

ORF_category_Gen: ORF annotation with respect to its genomic position .

NMD_candidate: TRUE or FALSE, depending on the presence of an additional exon-exon junction downstream the stop codon.

Distance_to_lastExEx: Distance (in nt) between the last exon-exon junction and the stop codon.

Author(s)

Lorenzo Calviello, <calviello.l.bio@gmail.com>

See Also

```
select_quantify_ORFs, annotate_splicing
```

annotate_splicing

Annotate splice features of detected ORFs

Description

This function detects usage of different exons and exonic boundaries of one ORF with respect to a reference ORF.

Usage

```
annotate_splicing(orf_gen, ref_cds)
```

Arguments

orf_gen Exon structure of a detected ORF ref_cds Exon structure of a reference ORF

Details

each exon is aligned to the closest one to match acceptor and donor sites, or to annotate missing exons. 5ss and 3ss indicate exon 5' and 3', respectively. CDS_spanning indicates retained intron; missing_CDS indicates no overlapping exon (missed or included); monoCDS indicates a single-exon ORF; firstCDS and lastCDS indicate first CDS exon or last CDS exon.

Value

Exon structure of detected ORF including possible missing exons from reference, together with a spl_type column including the annotation for each exon (e.g. alternative acceptors or donor).

Author(s)

Lorenzo Calviello, <calviello.1.bio@gmail.com>

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See Also

detect_translated_orfs, annotate_ORFs

calc_orf_pval

Collect ORF Ribo-seq statistics

Description

This function calculates statistics for the analysis of P_sites profiles for each ORF

Usage

```
calc_orf_pval(ORFs, P_sites_rle, P_sites_uniq_rle, P_sites_uniq_mm_rle,
  cutoff = 0.5, tapers = 24, bw = 12)
```

Arguments

ORFs Set of detected ORFs

P_sites_rle Rle signal of P_sites along the transcript

P_sites_uniq_rle
Rle signal of uniquely mapping P_sites along the transcript

P_sites_uniq_mm_rle
Rle signal of uniquely mapping P_sites with mismatches along the transcript

cutoff cutoff of average in-frame signal for each codon in the ORF. Defaults to .5

tapers Number of tapers to use in the multitaper analysis. Defaults to 24

Details

bw

Number of P_sites (uniquely mapping or all), frame percentage and multitaper test statistics are collected for each ORF. The parameter space for the multitaper analysis was explored in the RiboTaper paper.

time_bw parameter to use in the multitaper analysis. Defaults to 12

Value

Set of detected ORFs, including info about the possible longest ORF for that frame.

Author(s)

```
Lorenzo Calviello, <calviello.1.bio@gmail.com>
```

```
detect_translated_orfs, get_orfs, take_Fvals_spect
```

create_SaTAnn_html_report

Create an html report summarizing SaTAnn results

Description

This function creates an html report showing summary statistics for SaTAnn-detected ORFs.

Usage

```
create_SaTAnn_html_report(input_files, input_sample_names, output_file)
```

Arguments

Character vector containing input names. Must be of same length as input_files.

output_file String; full path to html report file.

Details

This function creates the html report visualizing final SaTAnn results.

Input are two lists of the same length:

- a) input_files: list of full paths to one or multiple input files (*SaTAnn_plots_RData files generated with plot_SaTAnn_results) and
- b) input_sample_names: list of corresponding names describing the file content (these are used as names in the report).

For the report, a RMarkdown file is rendered as html document, saved as output_file.

Value

The function saves the html report file with the file path output_file.

Author(s)

Lorenzo Calviello, <calviello.bio@gmail.com>

```
plot_SaTAnn_results, run_SaTAnn
```

detect_readthrough 7

| detect_readthrough Analyzed translation on possible readthrough regions (beta) |
|--|
|--|

Description

This function uses the multitaper method to look for readthrough translation

Usage

```
detect_readthrough(results_orf, P_sites, P_sites_uniq, P_sites_uniq_mm,
  genome_sequence, annotation, genetic_code_table, cutoff_fr_ave = 0.5,
  uniq_signal = F)
```

Arguments

results_orf Full list of detected ORFs, from select_quantify_ORFs and annotate_ORFs GRanges object with P_sites positions P_sites P_sites_uniq GRanges object with uniquely mapping P_sites positions P_sites_uniq_mm Rle signal of uniquely mapping P_sites with mismatches along the transcript genome_sequence BSgenome object annotation Rannot object containing annotation of CDS and transcript structures (see prepare_annotation_files) genetic_code_table GENETIC_CODE table to use cutoff_fr_ave cutoff parameter for the calc_orf_pval functions

Use only signal from uniquely mapping reads? Defaults to FALSE.

Details

uniq_signal

The function looks for stop-stop pairs after the stop codon of the detected ORF

Value

GRanges object with the set of translated readthrough regions

Author(s)

```
Lorenzo Calviello, <calviello.1.bio@gmail.com>
```

```
detect_translated_orfs, select_quantify_ORFs, annotate_ORFs, get_reathr_seq
```

```
detect_translated_orfs
```

Detect actively translated ORFs

Description

This function detects translated ORFs

Usage

```
detect_translated_orfs(selected_txs, genome_sequence, annotation, P_sites,
   P_sites_uniq, P_sites_uniq_mm, genomic_region, genetic_code,
   all_starts = T, nostarts = F, start_sel_cutoff = NA,
   start_sel_cutoff_ave = 0.5, cutoff_fr_ave = 0.5, uniq_signal = F)
```

Arguments

```
set of selected transcripts, output from select_txs
selected_txs
genome_sequence
                 BSgenome object
annotation
                 Rannot object containing annotation of CDS and transcript structures (see prepare_annotation_files)
                 GRanges object with P_sites positions
P_sites
P_sites_uniq
                 GRanges object with uniquely mapping P_sites positions
{\tt P\_sites\_uniq\_mm}
                 GRanges object with uniquely mapping (with mismatches) P_sites positions
genomic_region GRanges object with genomic coordinates of the genomic region analyzed
                 GENETIC_CODE table to use
genetic_code
all_starts
                 get_all_starts parameter for the get_orfs function
nostarts
                 Stop_Stop parameter for the get_orfs function
start_sel_cutoff
                 cutoff parameter for the select_start function
start_sel_cutoff_ave
                 cutoff_ave parameter for the select_start function
cutoff_fr_ave cutoff parameter for the calc_orf_pval functions
uniq_signal
                 Use only signal from uniquely mapping reads? Defaults to FALSE.
```

Details

A set of transcripts, together with genome sequence and Ribo-signal are analyzed to extract translated ORFs

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Value

A list with transcript coordinates, exonic coordinates and statistics for each ORF exonic bin and junction(from select_txs).

The value for each column is as follows:

ave_pct_fr: average percentage of in-frame reads for each codon in the ORF pct_fr: percentage of in-frame reads in the ORF ave_pct_fr: average percentage of in-frame reads for each codon in the ORF ave_pct_fr_st: average percentage of in-frame reads per each codon between the selected start codon and the next candidate one pct_fr_st: percentage of in-frame reads between the selected start codon and the next candidate one longest_ORF: GRanges coordinates for the longest ORF with the same stop codon pval: P-value for the multitaper F-test at 1/3 using the ORF P_sites profile pval_uniq: P-value for the multitaper F-test at 1/3 using the ORF P_sites profile (only uniquely mapping reads) P_sites_raw: Raw number of P_sites mapping to the ORF P_sites_raw_unique: Uniquely mapping P_sites mapping to the ORF ORF_id_tr: ORF id containing <tx_id>_<start>_<end> Protein: AAString sequence of the translated protein region: Genomic coordinates of the analyzed region gene_id: gene_id for the corresponding analyzed transcript gene_biotype: gene biotype for the corresponding analyzed transcript_id for the corresponding analyzed ORF transcript_biotype: transcript biotype for the corresponding analyzed ORF

Author(s)

Lorenzo Calviello, <calviello.1.bio@gmail.com>

See Also

```
select_txs, get_orfs, take_Fvals_spect, select_start, prepare_annotation_files
```

from_tx_togen

Map transcript coordinates to genomic coordinates

Description

This function uses the mapFromTranscripts function to switch between transcript and genomic coordinates

Usage

```
from_tx_togen(ORFs, exons, introns)
```

Arguments

| introns | intronic regions of the analyzed transcripts, as a GRangesList object |
|---------|---|
| exons | exonic regions of the analyzed transcripts, as a GRangesList object |
| ORFs | Set of detected ORFs from the calc_orf_pval function |

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Value

exonic coordinates for each ORF.

Author(s)

Lorenzo Calviello, <calviello.1.bio@gmail.com>

See Also

mapFromTranscripts

get_orfs

Find ATG-starting ORFs in a sequence

Description

This function loads the annotation created by the prepare_annotation_files function

Usage

```
get_orfs(tx_name, sequence, get_all_starts = T, Stop_Stop = F,
  scores = c(1, 0.5), genetic_code_table)
```

Arguments

tx_name transcript_id DNAString object containing the sequence of the transcript sequence

get_all_starts Output all possible start codons? Defaults to TRUE

Stop_Stop Find Stop-Stop pairs (no defined start codon)? Defaults to FALSE

Deprecated scores

genetic_code_table

GENETIC_CODE table to use

Value

GRanges object containing coordinates for the detected ORFs

Author(s)

```
Lorenzo Calviello, <calviello.l.bio@gmail.com>
```

```
detect_translated_orfs
```

get_ps_fromsplicemin 11

```
get_ps_fromsplicemin Offset spliced reads on minus strand
```

Description

This function calculates P-sites positions for spliced reads on the minus strand

Usage

```
get_ps_fromsplicemin(x, cutoff)
```

Arguments

x a GAlignments object with a cigar string cutoff number representing the offset value

Value

a GRanges object with offset reads

Author(s)

```
Lorenzo Calviello, <calviello.l.bio@gmail.com>
```

See Also

```
prepare_for_SaTAnn
```

```
{\tt get\_ps\_fromspliceplus} \ \ \textit{Offset spliced reads on plus strand}
```

Description

This function calculates P-sites positions for spliced reads on the plus strand

Usage

```
get_ps_fromspliceplus(x, cutoff)
```

Arguments

x a GAlignments object with a cigar string cutoff number representing the offset value

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Value

a GRanges object with offset reads

Author(s)

Lorenzo Calviello, <calviello.l.bio@gmail.com>

See Also

```
prepare_for_SaTAnn
```

get_reathr_seq

Extract possible readthrough sequences (beta)

Description

This function extracts readthrough regions for subsequent analysis

Usage

```
get_reathr_seq(tx_name, orf, sequence, genetic_code)
```

Arguments

tx_name transcript_id

orf transcript-level ORF coordinates

sequence DNAString object containing the sequence of the transcript

genetic_code GENETIC_CODE table to use

Details

The function looks for stop-stop pairs after the stop codon of the detected ORF

Value

GRanges object with the set of possible readthrough sequences

Author(s)

```
Lorenzo Calviello, <calviello.l.bio@gmail.com>
```

```
detect_translated_orfs, select_quantify_ORFs
```

load_annotation 13

load_annotation

Load genomic features and genome sequence

Description

This function loads the annotation created by the prepare_annotation_files function

Usage

```
load_annotation(path)
```

Arguments

path

Full path to the *Rannot R file in the annotation directory used in the prepare_annotation_files funct

Value

introduces a GTF_annotation object and a genome_seq object in the parent environment

Author(s)

Lorenzo Calviello, <calviello.l.bio@gmail.com>

See Also

```
prepare_annotation_files
```

plot_SaTAnn_results

Plot general statistics about SaTAnn results

Description

This function produces a series of plots and statistics about the set ORFs called by SaTAnn compared to the annotation. IMPORTANT: Use only on transcriptome-wide SaTAnn results. See run_SaTAnn

Usage

```
plot_SaTAnn_results(for_SaTAnn_file, SaTAnn_output_file, annotation_file,
  coverage_file_plus = NA, coverage_file_minus = NA,
  output_plots_path = NA, prefix = NA)
```

Arguments

for_SaTAnn_file

path to the "for_SaTAnn" file containing P_sites positions and junction reads

SaTAnn_output_file

Full path to the "_final_SaTAnn_results" RData object output by SaTAnn. See run_SaTAnn

annotation_file

coverage_file_plus

Full path to a Ribo-seq coverage (no P-sites but read coverage) bigwig file (plus strand), as the ones created by RiboseQC

coverage_file_minus

Full path to a Ribo-seq coverage (no P-sites but read coverage) bigwig file (minus strand), as the ones created by RiboseQC

output_plots_path

Full path to the directory where plots in .pdf format are stored.

prefix prefix appended to output filenames

Value

the function exports a RData object (*SaTAnn_plots_RData) containing data to produce all plots, and produces different QC plots in .pdf format. The plots created are as follows:

ORFs_found: Number of ORF categories detected per gene biotype.

ORFs_found_pct_tr: Distribution of ORF_pct_P_sites (ORFs_found_ORFs_pM: Distribution of ORFs_pM (ORFs per Million, similar to TPM) for different ORF categories and gene biotypes.

ORFs_found_len: Distribution of ORF length for different ORF categories and gene biotypes.

ORFs_genes: Number of detected ORFs per gene.

ORFs_genes_tpm: Gene level TPM values, plotted by number of ORFs detected.

ORFs_maxiso: Number of genes plotted against the percentages of gene translation of their most translated ORF.

ORFs_maxiso_tpm: Gene level TPM values, plotted against the percentages of gene translation of their most translated ORF.

Sel_txs_genes: Number of genes plotted against the number of selected transcripts.

Sel_txs_genes_tpm: Gene level TPM values, plotted against the number of selected transcripts.

Sel_txs_genes_pct: Percentages of annotated trascripts per gene, plotted against the number of selected transcripts.

Sel_txs_bins_juns: Percentages of covered exonic bins or junctions, using all annotated transcripts, coding transcripts only, or the set of selected transcripts.

Meta_splicing_coverage: Aggregate signal of Ribo-seq coverage and normalized ORF coverage across different splice sites combinations, with different mixtures of translated overlapping ORFs.

Author(s)

Lorenzo Calviello, <calviello.1.bio@gmail.com>

See Also

run_SaTAnn

```
prepare_annotation_files
```

Prepare comprehensive sets of annotated genomic features

Description

This function processes a gtf file and a twobit file (created using faToTwoBit from ucsc tools: http://hgdownload.soe.ucsc.edu/admin/exe/) to create a comprehensive set of genomic regions of interest in genomic and transcriptomic space (e.g. introns, UTRs, start/stop codons). In addition, by linking genome sequence and annotation, it extracts additional info, such as gene and transcript biotypes, genetic codes for different organelles, or chromosomes and transcripts lengths.

Usage

```
prepare_annotation_files(annotation_directory, twobit_file, gtf_file,
    scientific_name = "Homo.sapiens", annotation_name = "genc25",
    export_bed_tables_TxDb = T, forge_BSgenome = T, create_TxDb = T)
```

Arguments

annotation_directory

The target directory which will contain the output files

twobit_file Full path to the genome file in twobit format gtf_file Full path to the annotation file in GTF format

scientific_name

A name to give to the organism studied; must be two words separated by a ".",

defaults to Homo.sapiens

annotation_name

A name to give to annotation used; defaults to genc25

export_bed_tables_TxDb

Export coordinates and info about different genomic regions in the annotation_directory?

It defaults to TRUE

forge_BSgenome Forge and install a BSgenome package? It defaults to TRUE create_TxDb Create a TxDb object and a *Rannot object? It defaults to TRUE

Details

This function uses the makeTxDbFromGFF function to create a TxDb object and extract genomic regions and other info to a *Rannot R file; the mapToTranscripts and mapFromTranscripts functions are used to map features to genomic or transcript-level coordinates. GTF file mist contain "exon" and "CDS" lines, where each line contains "transcript_id" and "gene_id" values. Additional values such as "gene_biotype" or "gene_name" are also extracted. Regarding sequences, the twobit

file, together with input scientific and annotation names, is used to forge and install a BSgenome package using the forgeBSgenomeDataPkg function.

The resulting GTF_annotation object (obtained after running load_annotation) contains:

txs: annotated transcript boundaries.

txs_gene: GRangesList including transcript grouped by gene.

seqinfo: indicating chromosomes and chromosome lengths.

start_stop_codons: the set of annotated start and stop codon, with respective transcript and gene_ids. reprentative_mostcommon,reprentative_boundaries and reprentative_5len represent the most common start/stop codon, the most upstream/downstream start/stop codons and the start/stop codons residing on transcripts with the longest 5'UTRs

cds_txs: GRangesList including CDS grouped by transcript.

introns_txs: GRangesList including introns grouped by transcript.

cds_genes: GRangesList including CDS grouped by gene.

exons_txs: GRangesList including exons grouped by transcript.

exons_bins: the list of exonic bins with associated transcripts and genes.

junctions: the list of annotated splice junctions, with associated transcripts and genes.

genes: annotated genes coordinates.

threeutrs: collapsed set of 3'UTR regions, with correspinding gene_ids. This set does not overlap CDS region.

fiveutrs: collapsed set of 5'UTR regions, with correspinding gene_ids. This set does not overlap CDS region.

ncIsof: collapsed set of exonic regions of protein_coding genes, with correspinding gene_ids. This set does not overlap CDS region.

ncRNAs: collapsed set of exonic regions of non_coding genes, with correspinding gene_ids. This set does not overlap CDS region.

introns: collapsed set of intronic regions, with correspinding gene_ids. This set does not overlap exonic region.

intergenicRegions: set of intergenic regions, defined as regions with no annotated genes on either strand.

trann: DataFrame object including (when available) the mapping between gene_id, gene_name, gene_biotypes, transcript_id and transcript_biotypes.

cds_txs_coords: transcript-level coordinates of ORF boundaries, for each annotated coding transcript. Additional columns are the same as as for the start_stop_codons object.

genetic_codes: an object containing the list of genetic code ids used for each chromosome/organelle. see GENETIC CODE TABLE for more info.

genome_package: the name of the forged BSgenome package. Loaded with load_annotation function.

stop_in_gtf: stop codon, as defined in the annotation.

Value

a TxDb file and a *Rannot files are created in the specified annotation_directory. In addition, a BSgenome object is forged, installed, and linked to the *Rannot object

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Author(s)

Lorenzo Calviello, <calviello.1.bio@gmail.com>

See Also

load_annotation, forgeBSgenomeDataPkg, makeTxDbFromGFF, run_SaTAnn.

prepare_for_SaTAnn

Prepare the "for_SaTAnn" file

Description

Prepare the "for_SaTAnn" file

Usage

```
prepare_for_SaTAnn(annotation_file, bam_file,
  path_to_rl_cutoff_file = NA, chunk_size = 5e+06,
  path_to_P_sites_plus_bw = NA, path_to_P_sites_minus_bw = NA,
  path_to_P_sites_uniq_plus_bw = NA,
  path_to_P_sites_uniq_minus_bw = NA,
  path_to_P_sites_uniq_mm_plus_bw = NA,
  path_to_P_sites_uniq_mm_minus_bw = NA, dest_name = NA)
```

Arguments

```
annotation_file
```

Full path to the annotation file (*Rannot)

bam_file Full path to the bam file

path_to_rl_cutoff_file

path to the rl_cutoff_file file specifying in 3 columns the read lengths, cutoffs and compartments ("nucl" for standard chromosomes)

chunk_size the number of alignments to read at each iteration, defau

nk_size the number of alignments to read at each iteration, defaults to 5000000, increase when more RAM is available

path_to_P_sites_plus_bw

path to a bigwig file containing P_sites positions on the plus strand

path_to_P_sites_minus_bw

path to a bigwig file containing P_sites positions on the minus strand

path_to_P_sites_uniq_plus_bw

(Optional) path to a bigwig file containing uniquely mapping P_sites positions on the plus strand

path_to_P_sites_uniq_minus_bw

(Optional) path to a bigwig file containing uniquely mapping P_sites positions on the minus strand

run_SaTAnn

Details

This function uses a list of pre-determined read lengths, cutoffs and compartments to calculate P_sites positions.

Alternatively, bigwig files containing P_sites position for each strand can be specified. Optional bigwig files for uniquely mapping P_sites position (with and without mismatches) can be specified to obtain more statistics on the SaTAnn-identified ORFs

Author(s)

Lorenzo Calviello, <calviello.1.bio@gmail.com>

See Also

run_SaTAnn

run_SaTAnn

Run the SaTAnn pipeline

Description

This wrapper function runs the entire SaTAnn pipeline

Usage

```
run_SaTAnn(for_SaTAnn_file, annotation_file, n_cores,
   prefix = for_SaTAnn_file, gene_name = NA, gene_id = NA,
   genomic_region = NA, write_temp_files = T, write_GTF_file = T,
   write_protein_fasta = T, interactive = T,
   stn.orf_find.all_starts = T, stn.orf_find.nostarts = F,
   stn.orf_find.start_sel_cutoff = NA,
   stn.orf_find.cutoff_fr_ave = 0.5, stn.orf_quant.cutoff_cums = NA,
   stn.orf_quant.cutoff_pct = 2, stn.orf_quant.cutoff_P_sites = NA,
   unique_reads_only = F, canonical_start_only = T)
```

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Arguments

for_SaTAnn_file REQUIRED - path to the "for_SaTAnn" file containing P_sites positions and junction reads annotation_file REQUIRED - path to the *Rannot R file in the annotation directory used in the prepare_annotation_files function REQUIRED - number of cores to use n_cores prefix prefix to use for the output files. Defaults to same as for_SaTAnn_file (appends to its filename) character vector of gene names to analyze. gene_name gene_id character vector of gene ids to analyze genomic_region GRanges object with genomic regions to analyze write_temp_files write temporary files. Defaults to TRUE write_GTF_file write a GTF files with the ORF coordinates. Defaults to TRUE write_protein_fasta write a protein fasta file. Defaults to TRUE interactive should put R object in global environment? Defaults to TRUE stn.orf_find.all_starts orf_find.all_starts parameter for the SaTAnn function stn.orf_find.nostarts orf_find.nostarts parameter for the SaTAnn function stn.orf_find.start_sel_cutoff orf_find.start_sel_cutoff parameter for the SaTAnn function stn.orf_find.start_sel_cutoff_ave orf_find.start_sel_cutoff_ave parameter for the SaTAnn functio stn.orf_find.cutoff_fr_ave orf_find.cutoff_fr_ave parameter for the SaTAnn function stn.orf_quant.cutoff_cums orf_quant.cutoff_cums parameter for the SaTAnn function stn.orf_quant.cutoff_pct orf_quant.cutoff_pct parameter for the SaTAnn function stn.orf_quant.cutoff_P_sites orf_quant.cutoff_P_sites parameter for the SaTAnn function unique_reads_only Use only signal from uniquely mapping reads? Defaults to FALSE. canonical_start_only Use only the canonical start codon (no alternative initiation codons)? Defaults to TRUE.

Details

A set of transcripts, together with genome sequence and Ribo-signal are analyzed to extract translated ORFs

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Value

A set of output files containing transcript coordinates, exonic coordinates and annotation for each ORF, including optional GTF and protein fasta files.

The description for each list object is as follows:

tmp_SaTAnn_results: (Optional) RData object file containing the entire set of results for each genomic region.

final_SaTAnn_results: RData object file containing the final SaTAnn results, see SaTAnn. Protein_sequences.fasta: (Optional) Fasta file containing the set of translated proteins. Detected_ORFs.gtf: GTF file containing coordinates of the detected ORFs.

In addition, new columns are added in the ORFs_tx file:

ORFs_pM: number of P_sites for each ORF, divided by ORF length and summing up to a million (akin to TPM).

Author(s)

Lorenzo Calviello, <calviello.1.bio@gmail.com>

See Also

prepare_annotation_files, load_annotation, SaTAnn

SaTAnn

Detection, quantification and annotation of translated ORFs in a genomic region

Description

This function detects, quantifies and annotates actively translated ORF in a genomic region

Usage

```
SaTAnn(region, for_SaTAnn, genetic_code_region, orf_find.all_starts = T,
  orf_find.nostarts = F, orf_find.start_sel_cutoff = NA,
  orf_find.start_sel_cutoff_ave = 0.5, orf_find.cutoff_fr_ave = 0.5,
  orf_quant.cutoff_cums = NA, orf_quant.cutoff_pct = 2,
  orf_quant.cutoff_P_sites = NA, unique_reads = F)
```

Arguments

region GRanges object with genomic coordinates of the genomic region analyzed for_SaTAnn "for_SaTAnn" Robject containing P_sites positions and junction reads

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```
genetic_code_region
                GENETIC_CODE table to use
orf_find.all_starts
                 get_all_starts parameter for the detect_translated_orfs function
orf_find.nostarts
                 Stop_Stop parameter for the detect_translated_orfs function
orf_find.start_sel_cutoff
                 cutoff parameter for the detect_translated_orfs function
orf_find.start_sel_cutoff_ave
                cutoff_ave parameter for the detect_translated_orfs function
orf_find.cutoff_fr_ave
                cutoff parameter for the detect_translated_orfs function
orf_quant.cutoff_cums
                cutoff_cums parameter for the select_quantify_ORFs function
orf_quant.cutoff_pct
                cutoff_pct parameter for the select_quantify_ORFs function
orf_quant.cutoff_P_sites
                cutoff_P_sites parameter for the select_quantify_ORFs function
                Use only signal from uniquely mapping reads? Defaults to FALSE.
unique_reads
```

Details

A set of transcripts, together with genome sequence and Ribo-signal are analyzed to extract translated ORFs

Value

A list containing transcript coordinates, exonic coordinates and annotation for each ORF.

The description for each list object is as follows:

ORFs_tx: transcript coordinates of the detected ORFs.

ORFs_gen: genomic (exon) coordinates of the detected ORFs.

ORFs_feat: list of ORF features together with mapping reads and uniqueness.

ORFs_txs_feats: list of transcript features present in the genomic region, together with mapping reads and uniqueness.

ORFs_spl_feat_longest: splicing annotation for each ORF exon, with respect to the longest annotated coding transcript for each gene.

ORFs_spl_feat_maxORF: splicing annotation for each ORF exon, with respect to the most translated ORF in each gene.

selected_txs: character vector containing the transcript ids of the selected transcripts.

ORFs_readthroughs: (Beta) transcript coordinates of the detected ORFs readthroughs.

Author(s)

Lorenzo Calviello, <calviello.1.bio@gmail.com>

See Also

select_txs, detect_translated_orfs, select_quantify_ORFs, annotate_ORFs, detect_readthrough

```
select_quantify_ORFs Select and quantify ORF translation
```

Description

This function selects a subset of detected ORFs and quantifies their translation

Usage

```
select_quantify_ORFs(results_ORFs, P_sites, P_sites_uniq,
  cutoff_cums = NA, cutoff_pct = 2, cutoff_P_sites = NA,
  optimiz = FALSE, scaling = TRUE, uniq_signal = F)
```

Arguments

| results_ORFs | Full list of detected ORFs, from detect_translated_ORFs |
|----------------|---|
| P_sites | GRanges object with P_sites positions |
| P_sites_uniq | GRanges object with uniquely mapping P_sites positions |
| cutoff_cums | cutoff to select ORFs until $<$ x $>$ percentage of total gene translation. Defaults to 99 |
| cutoff_pct | minimum percentage of total gene translation for an ORF to be selected. Defaults to $\boldsymbol{1}$ |
| cutoff_P_sites | minimum number of P_sites assigned to the ORF to be selected. Defaults to 10 |
| optimiz | (Beta) should numerical optimization (minimizing distance between observed coverage and expected coverage) be used to quantify ORF translation? Defaults to FALSE |
| scaling | Additional scaling value taking into account total signal on the detected ORFs to adjust quantification estimates (recommended). Defaults to TRUE |
| uniq_signal | Use only signal from uniquely mapping reads? Defaults to FALSE. |
| | |

Details

ORFs are first selected using the same method as in the select_txs function, but using ORF features (ORF structures are treated as transcript structures).

Ribo-seq coverage (reads/length) on bins and junctions (set to a length of 60) is used to derive a scaling factor (0-1) for each ORF, which indicates how much of the ORF coverage can be assigned to such ORF (1 when no other ORF is present). When no unique features are present on an ORF, an adjusted scaling value is calculated subtracting coverage expected from a ORF with a unique feature. When no unique features are present on any ORF, scaling values are calculated assuming uniform coverage on each ORF.

ORFs are then further filtered to exclude lowly translated ORFs and quantification/selection is reiterated until no ORF is further filtered out. Percentage of total gene translation and length-adjusted

select_start 23

quantification estimates are produced. More details about the quantificatin procedure can be found in the SaTAnn manuscript.

Additional columns are added to the ORFs_tx object:

P_sites: P_sites_raw value from detect_translated_ORFs divided by the ORF scaling value.

 $ORF_pct_P_sites$: Percentage of gene translation output for the ORF, derived using P_sites values. $ORF_pct_P_sites_pN$: Percentage of gene translation output (adjusted by length) for the ORF, derived using P_sites values.

unique_features_reads: initial number of reads on each unique ORF feature. NA when no unique feature is present.

adj_unique_features_reads: final number of reads on each unique ORF feature after the ORF filtering/quantification procedure. NA when no unique feature is present.

scaling_factors: Set of 3 scaling factors assigned to the ORF using intial unique ORF features, after adjusting for the presence of ORFs with no unique features, and final scaling factor after correcting for total Ribo-seq coverage on the gene.

Value

modified results_ORFs object with the selected ORFs including quantification estimates.

Author(s)

Lorenzo Calviello, <calviello.l.bio@gmail.com>

See Also

detect_translated_orfs, select_txs

select_start

Select start codon

Description

This function selects the start codon for ORFs in the same transcript

Usage

```
select_start(ORFs, P_sites_rle, cutoff = NA, cutoff_ave = 0.5)
```

Arguments

ORFs Set of detected ORFs

P_sites_rle Rle signal of P_sites along the transcript

cutoff cutoff of total in-frame signal between start codons (sensitive to outliers). De-

faults to NA

cutoff_ave cutoff for frequency of in-frame codons between two start codons (less sensitive

to outliers). Defaults to .5

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Details

ORFs are divided based on stop codon and Ribo-seq signal between start codons is used to select one

When more than cutoff_ave fraction of codons is in-frame between two candidate start codons, the most upstream is selected.

Value

Set of detected ORFs, including info about the possible longest ORF for that frame.

Author(s)

Lorenzo Calviello, <calviello.1.bio@gmail.com>

See Also

```
detect_translated_orfs, get_orfs
```

select_txs

Select a subset of transcripts with Ribo-seq data

Description

This function flattens all annotated transcript structures and uses Ribo-seq to select a subset of transcripts.

Usage

```
select_txs(region, annotation, P_sites, P_sites_uniq, junction_counts,
  uniq_signal = F)
```

Arguments

region genomic region being analyzed

annotation Rannot object containing annotation of CDS and transcript structures (see prepare_annotation_files)

P_sites GRanges object with P_sites positions

P_sites_uniq GRanges object with uniquely mapping P_sites positions

junction_counts

GRanges object containing Ribo-seq counts on the set of annotated junctions

uniq_signal Use only signal from uniquely mapping reads? Defaults to FALSE.

Details

Features (bins and junctions) are divided into shared and unique features, and into with support and without support (with or without reads mapping). A set of logical rules filters out transcripts with internal features with no support and no unique features with reads. More specific details can be found in the SaTAnn manuscript.

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Value

GRanges object with the set of counts on each exonic bin and junctions, together with the list of selected transcripts

Author(s)

Lorenzo Calviello, <calviello.l.bio@gmail.com>

See Also

```
prepare_annotation_files
```

take_Fvals_spect

Extract output from multitaper analysis of a signal

Description

This function uses the multitaper tool to extract F-values and multitaper spectral coefficients

Usage

```
take_Fvals_spect(x, n_tapers, time_bw, slepians_values)
```

Arguments

x numeric signal to analyze

n_tapers n of tapers to use time_bw parameter

slepians_values

set of calculated slepian functions to use in the multitaper analysis

Details

Values reported correspond to the closest frequency to 1/3 (same parameters as in RiboTaper). Padding to a minimum length of 1024 is performed to increase spectral resolution.

Value

two numeric values representing the F-value for the multitaper test and its corresponding spectral coefficient at the closest frequency to 1/3

Author(s)

Lorenzo Calviello, <calviello.1.bio@gmail.com>

```
detect_translated_orfs, spec.mtm, dpss
```

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