## Package ‘raer’

**May 27, 2024**

<table>
<thead>
<tr>
<th><strong>Type</strong></th>
<th>Package</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Title</strong></td>
<td>RNA editing tools in R</td>
</tr>
<tr>
<td><strong>Version</strong></td>
<td>1.2.0</td>
</tr>
<tr>
<td><strong>Description</strong></td>
<td>Toolkit for identification and statistical testing of RNA editing signals from within R. Provides support for identifying sites from bulk-RNA and single cell RNA-seq datasets, and general methods for extraction of allelic read counts from alignment files. Facilitates annotation and exploratory analysis of editing signals using Bioconductor packages and resources.</td>
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<tr>
<td><strong>License</strong></td>
<td>MIT + file LICENSE</td>
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<tr>
<td><strong>Imports</strong></td>
<td>stats, methods, GenomicRanges, IRanges, Rsamtools, BSgenome, Biostrings, SummarizedExperiment, SingleCellExperiment, S4Vectors, GenomeInfoDb, GenomicAlignments, GenomicFeatures, BiocGenerics, BiocParallel, rtracklayer, Matrix, cli</td>
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<tr>
<td><strong>Suggests</strong></td>
<td>testthat (&gt;= 3.0.0), knitr, DESeq2, edgeR, limma, rmarkdown, BiocStyle, ComplexHeatmap, TxDb.Hsapiens.UCSC.hg38.knownGene, SNPloc.Hsapiens.dbSNP144.GRCh38, BSgenome.Hsapiens.NCBI.GRCh38, scater, scran, scuttle, AnnotationHub, covr, raerdata, txdbmaker</td>
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<td><a href="https://github.com/rnabioco/raer/issues">https://github.com/rnabioco/raer/issues</a></td>
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**biocViews** MultipleComparison, RNASeq, SingleCell, Sequencing, Coverage, Epitranscriptomics, FeatureExtraction, Annotation, Alignment

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Author  Kent Riemondy [aut, cre] (<https://orcid.org/0000-0003-0750-1273>),
Kristen Wells-Wrasman [aut] (<https://orcid.org/0000-0002-7466-8164>),
Ryan Sheridan [ctb] (<https://orcid.org/0000-0003-4012-3147>),
Jay Hesselberth [ctb] (<https://orcid.org/0000-0002-6299-179X>),
RNA Bioscience Initiative [cph, fnd]

Maintainer  Kent Riemondy <kent.riemondy@gmail.com>

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**annot_from_gr**

---

**Annotate sites using GRanges object**

**Description**

Utility function to map annotations from GRanges to rowData of SummarizedExperiment or to mcols of GRanges object. If multiple features overlap then they will be concatenated with the specified separator string.

**Usage**

```r
annot_from_gr(obj, gr, cols_to_map, RLE = TRUE, sep = ",", ...)```

**Arguments**

- `obj` RangedSummarizedExperiment or GRanges object
- `gr` GRanges with annotations to map to `obj`
- `cols_to_map` character vector of columns from GRanges to map to SummarizedExperiment. If the vector has names, the names will be the column names in the output.
- `RLE` If TRUE, columns added will returned as *S4Vectors::Rle()* vectors to reduce memory
- `sep` separator string, defaults to comma.
- `...` additional arguments to pass to *GenomicRanges::findOverlaps()*

**Value**

Either a SummarizedExperiment or GRanges object with additional annotations provided by the supplied GRanges object.

**Examples**

```r
library(SummarizedExperiment)
rse_adar_ifn <- mock_rse()
gr <- GRanges(rep(c("SSR3", "SPCS3"), c(5, 15)),
             IRanges(seq(1, 500, by = 25), width = 50), strand = "+")

gr$feature <- sample(1:100, size = 20)
gr$id <- sample(LETTERS, size = 20)

rse <- annot_from_gr(rse_adar_ifn, gr, c(feature_set = "feature", "id"))
rowData(rse)
```
**Annotate known SNP positions**

**Description**
This function will annotate a GRanges or the rowRanges of a SummarizedExperiment with SNPs from a SNP package.

**Usage**

```
annot_snps(obj, ...)
```

```r
## S3 method for class ‘GRanges’
annot_snps(
  obj,
  dbsnp,
  chrom = NULL,
  col_to_aggr = "RefSNP_id",
  drop = FALSE,
  genome = NULL,
  RLE = TRUE,
  ...
)
```

```r
## S3 method for class ‘SummarizedExperiment’
annot_snps(obj, ...)
```

**Arguments**
- `obj` GRanges or SummarizedExperiment object
- `...` For the generic, further arguments to pass to specific methods. Unused for now.
- `dbsnp` SNPloc package, see available packages from `BSgenome::available.SNPs()`.
- `chrom` only operate on a specified chromosome
- `col_to_aggr` column from SNPloc package to add to input. If multiple SNPs overlap these values will be concatenated as comma separated values.
- `drop` If TRUE, remove sites overlapping SNPs
- `genome` A BSgenome object, which if supplied, will be used to provide additional SNP_ref_allele and SNP_alt_alleles columns containing the reference and alt allele sequences, with respect to the positive strand. Additionally the SNP sequences will be checked against the allele at the site if a column named ALT is present in object. The strand of the site will be used to determine if the ALT allele needs to be complemented prior to comparing against the SNP db (which always returns sequences w.r.t the plus strand).
- `RLE` If TRUE, columns added will returned as S4Vectors::Rle() vectors to reduce memory usage.
calc_AEI

**Value**

Either a GRanges or SummarizedExperiment object with a new column added with information from col_to_aggr and optionally snp_ref_allele, snp_alt_alleles, and snp_matches_site annotations.

**See Also**

SNPlocs.Hsapiens.dbSNP144.GRCh38

**Examples**

```r
if (require(SNPlocs.Hsapiens.dbSNP144.GRCh38)) {
  gr <- GRanges(rep("22", 10),
    IRanges(
      seq(10510077, 10610077, by = 1000)
    )[1:10],
    width = 250
  ),
  strand = "+")
  genome(gr) <- "GRCh38.p2"
  annot_snps(gr, SNPlocs.Hsapiens.dbSNP144.GRCh38)
}
```

---

**Description**

The Adenosine Editing Index describes the magnitude of A-to-I editing in a sample. The index is a weighted average of editing events (G bases) observed at A positions. The vast majority A-to-I editing occurs in ALU elements in the human genome, and these regions have a high A-to-I editing signal compared to other regions such as coding exons. This function will perform pileup at specified repeat regions and return a summary AEI metric.

**Usage**

```r
calc_AEI(
  bamfiles, 
  fasta, 
  alu_ranges = NULL, 
  txdb = NULL, 
  snp_db = NULL, 
  param = FilterParam(), 
  BPPARAM = SerialParam(), 
  verbose = FALSE
)
```
Arguments

- **bamfiles**: character vector of paths to indexed bam files. If a named character vector is supplied the names will be used in the output.
- **fasta**: fasta filename
- **alu_ranges**: GRanges with regions to query for calculating the AEI, typically ALU repeats.
- **txdb**: A TxDb object, if supplied, will be used to subset the alu_ranges to those found overlapping genes. Alternatively a GRanges object with gene coordinates. If the library_type, specified by FilterParam, is unstranded then the TxDb will be used to correct the strandness relative to the reference and is a required parameter.
- **snp_db**: either a SNPlocs, GPos, or GRanges object. If supplied, will be used to exclude polymorphic positions prior to calculating the AEI. If calc_AEI() will be used many times, one will save time by first identifying SNPs that overlap the supplied alu_ranges, and passing these as a GRanges to snp_db rather than supplying all known SNPs (see get_overlapping_snps()).
- **param**: object of class FilterParam() which specify various filters to apply to reads and sites during pileup.
- **BPPARAM**: A BiocParallelParam object for specifying parallel options for operating over chromosomes.
- **verbose**: report progress on each chromosome?

Value

A named list containing:

- **AEI**: a matrix of AEI index values computed for all allelic combinations, one row for each supplied bam file.
- **AEI_per_chrom**: a data.frame containing values computed for each chromosome

References


Examples

```r
suppressPackageStartupMessages(library(Rsamtools))

bamfn <- raer_example("SRR5564269_Aligned.sortedByCoord.out.md.bam")
bam2fn <- raer_example("SRR5564277_Aligned.sortedByCoord.out.md.bam")
bams <- c(bamfn, bam2fn)
names(bams) <- c("ADAR1KO", "WT")

fafn <- raer_example("human.fasta")
mock_alu_ranges <- scanFaIndex(fafn)
```
calc_confidence

res <- calc_AEI(bams, fafn, mock_alu_ranges)
res$AEI

calc_confidence  
*Calculate confidence score for observing editing*

**Description**

Calculate a confidence score based on a Bayesian inverse probability model as described by Washburn et al. Cell Reports. 2015, and implemented in the SAILOR pipeline.

**Usage**

```r
calc_confidence(
    se,
    edit_to = "G",
    edit_from = "A",
    per_sample = FALSE,
    exp_fraction = 0.01,
    alpha = 0L,
    beta = 0L
)
```

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>se</td>
<td>SummarizedExperiment::SummarizedExperiment containing editing sites</td>
</tr>
<tr>
<td>edit_to</td>
<td>edited base</td>
</tr>
<tr>
<td>edit_from</td>
<td>non-edited base</td>
</tr>
<tr>
<td>per_sample</td>
<td>if TRUE, calculate confidence per sample, otherwise edited and non-edited counts will be summed across all samples.</td>
</tr>
<tr>
<td>exp_fraction</td>
<td>Numeric value between 0 and 1, specifying the expected error rate</td>
</tr>
<tr>
<td>alpha</td>
<td>Pseudo-count to add to non-edited base counts</td>
</tr>
<tr>
<td>beta</td>
<td>Pseudo-count to add to edited base counts</td>
</tr>
</tbody>
</table>

**Value**

SummarizedExperiment::SummarizedExperiment with either a new assay or rowData column named "confidence" depending on whether confidence is calculated per_sample.

**References**


SAILOR pipeline: https://github.com/YeoLab/sailor
Examples

```r
rse_adar_ifn <- mock_rse()
calc_confidence(rse_adar_ifn)
calc_confidence(rse_adar_ifn, per_sample = TRUE)
```

calc_edit_frequency  
*Adds editing frequencies*

Description

Adds editing frequencies to an existing *RangedSummarizedExperiment* object (created by `pileup_sites()`). The *RangedSummarizedExperiment* with a new assay for editing frequencies for each site (`edit_freq`), depth of coverage computed using the indicated edited nucleotides (`depth`) and new `colData` columns with the number of edited sites (`n_sites`) and the fraction of edits (`edit_idx`) is returned.

Usage

```r
calc_edit_frequency(
  rse,
  edit_from = "A",
  edit_to = "G",
  drop = FALSE,
  replace_na = TRUE,
  edit_frequency = 0,
  min_count = 1
)
```

Arguments

- `rse`: A *RangedSummarizedExperiment* object created by `pileup_sites()`
- `edit_from`: This should correspond to a nucleotide or assay (A, C, G, T, Ref, or Alt) you expect in the reference. Ex. for A to I editing events, this would be A.
- `edit_to`: This should correspond to a nucleotide or assay (A, C, G, T, Ref, or Alt) you expect in the editing site. Ex. for A to I editing events, this would be G.
- `drop`: If TRUE, the *RangedSummarizedExperiment* returned will only retain sites matching the specified `edit_from` and `edit_to` bases.
- `replace_na`: If TRUE, NA and NaN editing frequencies will be coerced to 0.
- `edit_frequency`: The edit frequency cutoff used when calculating the number of sites. Set to 0 to require any non-zero editing frequency. The number of sites is stored as `n_sites` in the `colData`.
- `min_count`: The minimum number of reads required when enumerating number of editing sites detected.
**calc_scAEI**

**Value**

*RangedSummarizedExperiment* supplemented with *edit_freq* and *depth* assay.

**Examples**

```r
library(SummarizedExperiment)

rse_adar_ifn <- mock_rse()

rse <- calc_edit_frequency(rse_adar_ifn)

assay(rse, "edit_freq")[1:5,]
```

---

**calc_scAEI**

*Calculate the Adenosine Editing Index (AEI) in single cells*

**Description**

The Adenosine Editing Index describes the magnitude of A-to-I editing in a sample. The index is a weighted average of editing events (G bases) observed at A positions. The vast majority A-to-I editing occurs in ALU elements in the human genome, and these regions have a high A-to-I editing signal compared to other regions such as coding exons. This function will examine enumerate edited and non-edited base counts at the supplied sites and return summary AEI metric per cell. Potential editing sites within repeat regions can be generated using *get_scAEI_sites()*.

**Usage**

```r
calc_scAEI(
  bamfiles,
  sites,
  cell_barcodes,
  param = FilterParam(),
  edit_from = "A",
  edit_to = "G",
  output_dir = NULL,
  return_sce = FALSE,
  ...
)

get_scAEI_sites(fasta, genes, alus, edit_from = "A", edit_to = "G")
```

**Arguments**

- **bamfiles**: a path to a BAM file (for 10x libraries), or a vector of paths to BAM files (smart-seq2). Can be supplied as a character vector, *BamFile*, or *BamFileList*.
- **sites**: a GRanges object produced by *get_scAEI_sites()* containing sites to process.
- **cell_barcodes**: A character vector of single cell barcodes to process. If processing multiple BAM files (e.g. smart-seq-2), provide a character vector of unique identifiers for each input BAM, to name each BAM file in the output files.
param: object of class `FilterParam()` which specify various filters to apply to reads and sites during pileup.

edit_from: This should correspond to the base (A, C, G, T) you expect in the reference. Ex. for A to I editing events, this would be A.

edit_to: This should correspond to the base (A, C, G, T) you expect in an edited site. Ex. for A to I editing events, this would be G.

output_dir: Output directory for nRef and nAlt sparseMatrix files. If NULL, a temporary directory will be used.

return_sce: if TRUE, data is returned as a SingleCellExperiment, if FALSE a DataFrame containing computed AEI values will be returned.

fasta: Path to a genome fasta file

genes: A GRanges object with gene coordinates. Alternatively a TxDb object, which if supplied, will be used extract gene coordinates.

alus: GRanges with repeat regions to query for calculating the AEI, typically ALU repeats. The strand of the supplied intervals will be ignored for defining repeat regions.

Value

A DataFrame containing computed AEI values, count of editing events (n_alt), and count of reference events (n_ref) per cell. If return_sce is TRUE, then a SingleCellExperiment is returned with the AEI values stored in the colData.

References


Examples

```r
suppressPackageStartupMessages(library(Rsamtools))
library(GenomicRanges)

bam_fn <- raer_example("5k_neuron_mouse_possort.bam")
bai <- indexBam(bam_fn)

# cell barcodes to query
cbs <- c("TGTTTGTTCCATCCGT-1", "CAACCAACATAATCGC-1", "TGGAACTCAAGCTGTT-1")

# genes used to infer transcribed strand
genes_gr <- GRanges(c(
  "2:100-400:-",
  "2:500-605:-",
  "2:600-680:+"
))
```
# alu intervals
alus_gr <- GRanges(c(
  "2:110-380",
  "2:510-600",
  "2:610-670"
))

# genome fasta file, used to find A bases
fa_fn <- raer_example("mouse_tiny.fasta")

# get positions of potential A -> G changes in alus
sites <- get_scAEI_sites(fa_fn, genes_gr, alus_gr)

fp <- FilterParam( 
  library_type = "fr-second-strand",
  min_mapq = 255
 ) 

calc_scAEI(bam_fn, sites, cbs, fp)

---

correct_strand  
Apply strand correction using gene annotations

Description

Gene annotations are used to infer the likely strand of editing sites. This function will operate on unstranded datasets which have been processed using "unstranded" library type which reports variants with respect to the + strand for all sites. The strand of the editing site will be assigned the strand of overlapping features in the genes_gr object. Sites with no-overlap, or overlapping features with conflicting strands (+ and -) will be removed.

Usage

correct_strand(rse, genes_gr)

Arguments

rse  
RangedSummarizedExperiment object containing editing sites processed with "unstranded" setting

genes_gr  
GRanges object containing reference features to annotate the strand of the editing sites.

Value

RangedSummarizedExperiment object containing pileup assays, with strand corrected based on supplied genomic intervals.
**Examples**

```r
suppressPackageStartupMessages(library("GenomicRanges"))

bamfn <- raer_example("SRR5564269_Aligned.sortedByCoord.out.md.bam")
fafn <- raer_example("human.fasta")
fp <- FilterParam(library_type = "unstranded")
rse <- pileup_sites(bamfn, fafn, param = fp)

genes <- GRanges(c(
  "DHFR:200-400:+",
  "SPCS3:100-200:-",
  "SSR3:3-10:-",
  "SSR3:6-12:+"
))

correct_strand(rse, genes)
```

---

**filter_clustered_variants**

*Filter out clustered sequence variants*

**Description**

Sequence variants of multiple allele types (e.g., A -> G, A -> C) proximal to a putative editing site can be indicative of a region prone to mis-alignment artifacts. Sites will be removed if variants of multiple allele types are present within a given distance in genomic or transcriptome coordinate space.

**Usage**

```r
filter_clustered_variants(
  rse,
  txdb,
  regions = c("transcript", "genome"),
  variant_dist = 100
)
```

**Arguments**

- **rse**: SummarizedExperiment::SummarizedExperiment containing editing sites
- **txdb**: GenomicFeatures::TxDb
- **regions**: One of transcript or genome, specifying the coordinate system for calculating distances between variants.
- **variant_dist**: distance in nucleotides for determining clustered variants
**filter_multiallelic**

**Value**

SummarizedExperiment::SummarizedExperiment with sites removed from object dependent on filtering applied.

**See Also**

Other se-filters: `filter_multiallelic()`, `filter_splice_variants()`

**Examples**

```r
if(require("txdbmaker")){
  rse_adar_ifn <- mock_rse()
  rse <- rse_adar_ifn[seqnames(rse_adar_ifn) == "SPCS3"]

  # mock up a txdb with genes
  gr <- GRanges(c(
    "SPCS3:100-120:-",
    "SPCS3:325-350:-"
  ))
  gr$source <- "raer"
  gr$type <- "exon"
  gr$source <- NA
  gr$phase <- NA
  gr$gene_id <- c(1, 2)
  gr$transcript_id <- c("1.1", "2.1")
  txdb <- txdbmaker::makeTxDbFromGRanges(gr)

  rse <- filter_multiallelic(rse)
  filter_clustered_variants(rse, txdb, variant_dist = 10)
}
```

---

**filter_multiallelic**  Filter out multi-allelic sites

**Description**

Remove sites with multiple variant bases from a SummarizedExperiment. `rowData()` gains a new column, `ALT`, that contains the variant allele detected at each site.

**Usage**

`filter_multiallelic(se)`

**Arguments**

- `se`  SummarizedExperiment::SummarizedExperiment
filter_splice_variants

Value

SummarizedExperiment::SummarizedExperiment with multiallelic sites removed. A new column, ALT will be added to rowData() indicating the single allele present at the site.

See Also

Other se-filters: filter_clustered_variants(), filter_splice_variants()

Examples

rse_adar_ifn <- mock_rse()
filter_multiallelic(rse_adar_ifn)

filter_splice_variants

Filter out sites near splice sites

Description

Remove editing sites found in regions proximal to annotated splice junctions.

Usage

filter_splice_variants(rse, txdb, splice_site_dist = 4, ignore.strand = FALSE)

Arguments

rse SummarizedExperiment::SummarizedExperiment with editing sites
txdb GenomicFeatures::TxDb
splice_site_dist distance to splice site
ignore.strand if TRUE, ignore strand when comparing editing sites to splice sites

Value

SummarizedExperiment::SummarizedExperiment with sites adjacent to splice sites removed.

See Also

Other se-filters: filter_clustered_variants(), filter_multiallelic()
**find_de_sites**

Perform differential editing

**Description**

Use edgeR or DESeq2 to perform differential editing analysis. This will work for designs that have 1 treatment and 1 control group. For more complex designs, we suggest you perform your own modeling.

**Usage**

```r
find_de_sites(
  deobj,
  test = c("edgeR", "DESeq2"),
  sample_col = "sample",
  condition_col = "condition",
  condition_control = NULL,
  condition_treatment = NULL
)
```

**Arguments**

- `deobj` A `RangedSummarizedExperiment` object prepared for differential editing analysis by `make_de_object()`
find_de_sites

- **test**: Indicate if edgeR or DESeq2 should be run.
- **sample_col**: The name of the column from colData(deobj) that contains your sample information. Default is sample. If you do not have a column named "sample", you must provide the appropriate sample column.
- **condition_col**: The name of the column from colData(deobj) that contains your treatment information. Default is condition. If you do not have a column named "condition", you must provide the appropriate condition column.
- **condition_control**: The name of the control condition. This must be a variable in your condition_col of colData(deobj). No default provided.
- **condition_treatment**: The name of the treatment condition. This must be a variable in your condition_col of colData(deobj).

**Value**

A named list:

- **de_obj**: The edgeR or deseq object used for differential editing analysis.
- **results_full**: Unfiltered differential editing results.
- **sig_results**: Filtered differential editing (FDR < 0.05).
- **model_matrix**: The model matrix used for generating DE results.

**Examples**

```r
library(SummarizedExperiment)
bamfn <- raer_example("SRR5564269_Aligned.sortedByCoord.out.md.bam")
bam2fn <- raer_example("SRR5564277_Aligned.sortedByCoord.out.md.bam")
fafn <- raer_example("human.fasta")

bams <- rep(c(bamfn, bam2fn), each = 3)
sample_ids <- paste0(rep(c("KO", "WT"), each = 3), 1:3)
names(bams) <- sample_ids

fp <- FilterParam(only_keep_variants = TRUE)
rse <- pileup_sites(bams, fafn, param = fp)
rse$condition <- substr(rse$sample, 1, 2)

rse <- calc_edit_frequency(rse)
dse <- make_de_object(rse)
res <- find_de_sites(dse,
  condition_control = "WT",
  condition_treatment = "KO"
)
res$sig_results[1:3,]
```
find_mispriming_sites

Description

OligoT will prime at A-rich regions in an RNA. Reverse transcription from these internal priming sites will install an oligoT sequence at the 3’ end of the cDNA. Sequence variants within these internal priming sites are enriched for variants converting the genomic sequence to the A encoded by the oligoT primer. Trimming poly(A) from the 3’ ends of reads reduces but does not eliminate these signals.

This function will identify regions that are enriched for mispriming events. Reads that were trimmed to remove poly(A) (encoded in the pa tag by 10x Genomics) are identified. The aligned 3’ positions of these reads are counted, and sites passing thresholds (at least 2 reads) are retained as possible sites of mispriming. By default regions 5 bases upstream and 20 bases downstream of these putative mispriming sites are returned.

Usage

```r
find_mispriming_sites(
  bamfile, 
  fasta, 
  pos_5p = 5, 
  pos_3p = 20, 
  min_reads = 2, 
  tag = "pa", 
  tag_values = 3:300, 
  n_reads_per_chunk = 1e+06,  
  verbose = TRUE
)
```

Arguments

- **bamfile**: path to bamfile
- **fasta**: path to fasta file
- **pos_5p**: distance 5’ of mispriming site to define mispriming region
- **pos_3p**: distance 3’ of mispriming site to define mispriming region
- **min_reads**: minimum required number of reads at a mispriming site
- **tag**: bam tag containing number of poly(A) bases trimmed
- **tag_values**: range of values required for read to be considered
- **n_reads_per_chunk**: number of reads to process in memory, see `Rsamtools::BamFile()`
- **verbose**: if true report progress
Value

A GenomicsRanges containing regions enriched for putative mispriming events. The n_reads column specifies the number of polyA trimmed reads overlapping the mispriming region. mean_pal indicates the mean length of polyA sequence trimmed from reads overlapping the region. The n_regions column specifies the number overlapping independent regions found in each chunk (dictated by n_reads_per_chunk). The A_freq column indicates the frequency of A bases within the region.

Examples

```r
bam_fn <- raer_example("5k_neuron_mouse_possort.bam")
fa_fn <- raer_example("mouse_tiny.fasta")
find_mispriming_sites(bam_fn, fa_fn)
```

find_scde_sites

Identify sites with differential editing between cells in single cell datasets

Description

Compare editing frequencies between clusters or celltypes. REF and ALT counts from each cluster are pooled to create pseudobulk estimates. Each pair of clusters are compared using fisher exact tests. Statistics are aggregated across each pairwise comparison using scran::combineMarkers.

Usage

```r
find_scde_sites(sce, group, rowData = FALSE, BPPARAM = SerialParam(), ...)
```

Arguments

- **sce** SingleCellExperiment object with nRef and nAlt assays.
- **group** column name from colData used to define groups to compare.
- **rowData** if TRUE, rowData from the input SingleCellExperiment will be included in the output DataFrames
- **BPPARAM** BiocParallel backend for control how parallel computations are performed.
- **...** Additional arguments passed to scran::combineMarkers

Value

A named list of DataFrames containing results for each cluster specified by group. The difference in editing frequencies between cluster pairs are denoted as dEF. See scran::combineMarkers for a description of additional output fields.
get_overlapping_snps

Retrieve SNPs overlapping intervals

Description
This function will find SNPs overlapping supplied intervals using a SNPlocs package. The SNPs can be returned in memory (as GPos objects) or written to disk as a bed-file (optionally compressed).

Usage
get_overlapping_snps(gr, snpDb, output_file = NULL)

Arguments

- **gr**: Intervals to query
- **snpDb**: A reference of a SNPlocs database
- **output_file**: A path to an output file. If supplied the file can be optionally compressed by including a ".gz" suffix. If not supplied, SNPs will be returned as a GenomicRanges::GPos object

Value
GPos object containing SNPs overlapping supplied genomic intervals
get_splice_sites

Examples

```r
if (require(SNPlocs.Hsapiens.dbSNP144.GRCh38)) {
  gr <- GRanges(rep("22", 10),
                IRanges(seq(10510077, 10610077, by = 1000)[1:10], width = 250),
                strand = "+")
  get_overlapping_snps(gr, SNPlocs.Hsapiens.dbSNP144.GRCh38)
}
```

get_splice_sites

Extract regions surrounding splice sites

Description

Extract intervals at splice sites and their adjacent regions.

Usage

```r
get_splice_sites(txdb, slop = 4)
```

Arguments

- `txdb` GenomicFeatures::TxDb
- `slop` The number of bases upstream and downstream of splice site to extract

Value

GenomicRanges::GRanges containing positions of splice sites, with flanking bases.

Examples

```r
if (require(TxDb.Hsapiens.UCSC.hg38.knownGene)) {
  txdb <- TxDb.Hsapiens.UCSC.hg38.knownGene
  res <- get_splice_sites(txdb)
  res[1:5]
}
```
make_de_object  

Make summarized experiment object for differential editing analysis

Description

Generates a `RangedSummarizedExperiment` object for use with edgeR or DESeq2. Will generate a counts assay with a matrix formatted with 2 columns per sample, representing the reference and editing allele counts.

Usage

```r
make_de_object(
  rse,
  edit_from = "A",
  edit_to = "G",
  min_prop = 0,
  max_prop = 1,
  min_samples = 1
)
```

Arguments

- `rse` A `RangedSummarizedExperiment` object
- `edit_from` This should correspond to a nucleotide or assay (A, C, G, T, Ref, or Alt) you expect in the reference. Ex. for A to I editing events, this would be A.
- `edit_to` This should correspond to a nucleotide or assay (A, C, G, T, Ref, or Alt) you expect in the editing site. Ex. for A to I editing events, this would be G.
- `min_prop` The minimum required proportion of reads edited at a site. At least `min_samples` need to pass this to keep the site.
- `max_prop` The maximum allowable proportion of reads edited at a site. At least `min_samples` need to pass this to keep the site.
- `min_samples` The minimum number of samples passing the `min_prop` and `max_prop` cutoffs to keep a site.

Value

`RangedSummarizedExperiment` for use with edgeR or DESeq2. Contains a counts assay with a matrix formatted with 2 columns per sample (ref and alt counts).

Examples

```r
library(SummarizedExperiment)
res_adar_ifn <- mock_rse()
res <- calc_edit_frequency(res_adar_ifn)
dse <- make_de_object(res, min_samples = 1)
assay(dse, "counts")[1:5,]
dse
```
mock_rse

Generate a small RangedSummarizedExperiment object for tests and examples

Description

A RangedSummarizedExperiment containing a subset of data from an RNA-seq experiment to measure the effects of IFN treatment of cell lines with wild-type or ADAR1-KO.

Usage

mock_rse()

Value

RangedSummarizedExperiment populated with pileup data

Source


References


Examples

mock_rse()

pileup_cells

Generate base counts per cell

Description

This function processes scRNA-seq library to enumerate base counts for Reference (unedited) or Alternate (edited) bases at specified sites in single cells. pileup_cells can process droplet scRNA-seq libraries, from a BAM file containing a cell-barcode and UMI, or well-based libraries that do not contain cell-barcodes.

The sites parameter specifies sites to quantify. This must be a GRanges object with 1 base intervals, a strand (+ or -), and supplemented with metadata columns named REF and ALT containing the reference and alternate base to query. See examples for the required format.

At each site, bases from overlapping reads will be examined, and counts of each ref and alt base enumerated for each cell-barcode present. A single base will be counted once for each UMI sequence present in each cell.
pileup_cells

Usage

pileup_cells(
  bamfiles,
  sites,
  cell_barcodes,
  output_directory,
  chroms = NULL,
  umi_tag = "UB",
  cb_tag = "CB",
  param = FilterParam(),
  BPPARAM = SerialParam(),
  return_sce = TRUE,
  verbose = FALSE
)

Arguments

bamfiles a path to a BAM file (for droplet scRNA-seq), or a vector of paths to BAM files (Smart-seq2). Can be supplied as a character vector, BamFile, or BamFileList.
sites a GRanges object containing sites to process. See examples for valid formatting.
cell_barcodes A character vector of single cell barcodes to process. If processing multiple BAM files (e.g. Smart-seq2), provide a character vector of unique identifiers for each input BAM, to name each BAM file in the output files.
output_directory Output directory for output matrix files. The directory will be generated if it doesn’t exist.
chroms A character vector of chromosomes to process. If supplied, only sites present in the listed chromosomes will be processed
umi_tag tag in BAM containing the UMI sequence
cb_tag tag in BAM containing the cell-barcode sequence
param object of class FilterParam() which specify various filters to apply to reads and sites during pileup. Note that the min_depth and min_variant_reads parameters if set > 0 specify the number of reads from any cell required in order to report a site. E.g. if min_variant_reads is set to 2, then at least 2 reads (from any cell) must have a variant in order to report the site. Setting min_depth and min_variant_reads to 0 reports all sites present in the sites object. The following options are not enabled for pileup_cells(): max_mismatch_type, homopolymer_len, and min_allelic_freq.
BPPARAM BiocParallel instance. Parallel computation occurs across chromosomes.
return_sce if TRUE, data is returned as a SingleCellExperiment, if FALSE a character vector of the output files, specified by outfile_prefix, will be returned.
verbose Display messages
Value

Returns either a SingleCellExperiment or character vector of paths to the sparseMatrix files produced. The SingleCellExperiment object is populated with two assays, nRef and nAlt, which represent base counts for the reference and alternate alleles. The rowRanges() will contain the genomic interval for each site, along with REF and ALT columns. The rownames will be populated with the format site_[seqnames]_[position(1-based)]_[strand]_[allele], with strand being encoded as 1 = +, 2 = -, and 3 = *, and allele being REF + ALT.

If return_sce is FALSE then a character vector of paths to the sparseMatrix files (barcodes.txt.gz, sites.txt.gz, counts.mtx.gz), will be returned. These files can be imported using read_sparray().

See Also

Other pileup: pileup_sites()

Examples

library(Rsamtools)
library(GenomicRanges)

bam_fn <- raer_example("5k_neuron_mouse_possort.bam")

gr <- GRanges(c("2:579:-", "2:625:-", "2:645:-", "2:589:-", "2:601:-"))
gr$REF <- c(rep("A", 4), "T")
gr$ALT <- c(rep("G", 4), "C")

cbs <- unique(scanBam(bam_fn, param = ScanBamParam(tag = "CB"))[[1]]$tag$CB)
cbs <- na.omit(cbs)

outdir <- tempdir()
bai <- indexBam(bam_fn)

fp <- FilterParam(library_type = "fr-second-strand")
sce <- pileup_cells(bam_fn, gr, cbs, outdir, param = fp)
sce

# example of processing multiple Smart-seq2 style libraries

many_small_bams <- rep(bam_fn, 10)
bam_ids <- LETTERS[1:10]

# for unstranded libraries, sites and alleles should be provided on + strand
gr$REF <- c(rep("T", 4), "A")
gr$ALT <- c(rep("C", 4), "G")

fp <- FilterParam(
    library_type = "unstranded",
    remove_overlaps = TRUE
)

sce <- pileup_cells(many_small_bams, sites = gr,
Description
This function uses a pileup routine to examine numerate base counts from alignments at specified sites, regions, or across all read alignments, from one or more BAM files. Alignment and site filtering options are controlled by the FilterParam class. A RangedSummarizedExperiment object is returned, populated with base count statistics for each supplied BAM file.

Usage
pileup_sites(
  bamfiles,
  fasta,
  sites = NULL,
  region = NULL,
  chroms = NULL,
  param = FilterParam(),
  BPPARAM = SerialParam(),
  umi_tag = NULL,
  verbose = FALSE
)

FilterParam(
  max_depth = 10000,
  min_depth = 1L,
  min_base_quality = 20L,
  min_mapq = 0L,
  library_type = "fr-first-strand",
  bam_flags = NULL,
  only_keep_variants = FALSE,
  trim_5p = 0L,
  trim_3p = 0L,
  ftrim_5p = 0,
  ftrim_3p = 0,
  indel_dist = 0L,
splice_dist = 0L,
min_splice_overhang = 0L,
homopolymer_len = 0L,
max_mismatch_type = c(0L, 0L),
read_bqual = c(0, 0),
min_variant_reads = 0L,
min_allelic_freq = 0,
report_multiallelic = TRUE,
remove_overlaps = TRUE
)

Arguments

bamfiles a character vector, BamFile or BamFileList indicating 1 or more BAM files to process. If named, the names will be included in the colData of the RangedSummarizedExperiment as a sample column, otherwise the names will be taken from the basename of the BAM file.

fasta path to genome fasta file used for read alignment. Can be provided in compressed gzip or bgzip format.
sites a GRanges object containing regions or sites to process.
region samtools region query string (i.e. chr1:100-1000). Can be combined with sites, in which case sites will be filtered to keep only sites within the region.
chroms chromosomes to process, provided as a character vector. Not to be used with the region parameter.
param object of class FilterParam() which specify various filters to apply to reads and sites during pileup.

BPPARAM A BiocParallel class to control parallel execution. Parallel processing occurs per chromosome and is disabled when run on a single region.

umi_tag The BAM tag containing a UMI sequence. If supplied, multiple reads with the same UMI sequence will only be counted once per position.
verbose if TRUE, then report progress and warnings.
max_depth maximum read depth considered at each site
min_depth min read depth needed to report site
min_base_quality min base quality score to consider read for pileup
min_mapq minimum required MAPQ score. Values for each input BAM file can be provided as a vector.
library_type read orientation, one of fr-first-strand, fr-second-strand, and unstranded. Unstranded library type will be reported with variants w.r.t the + strand. Values for each input BAM file can be provided as a vector.
bam_flags bam flags to filter or keep, use Rsamtools::scanBamFlag() to generate.
only_keep_variants if TRUE, then only variant sites will be reported (FALSE by default). Values for each input BAM file can be provided as a vector.
trim_5p  Bases to trim from 5’ end of read alignments
trim_3p  Bases to trim from 3’ end of read alignments
ftrim_5p  Fraction of bases to trim from 5’ end of read alignments
ftrim_3p  Fraction of bases to trim from 3’ end of read alignments
indel_dist  Exclude read if site occurs within given distance from indel event in the read
splice_dist  Exclude read if site occurs within given distance from splicing event in the read
min_splice_overhang  Exclude read if site is located adjacent to splice site with an overhang less than given length.
homopolymer_len  Exclude site if occurs within homopolymer of given length
max_mismatch_type  Exclude read if it has X different mismatch types (e.g. A-to-G, G-to-C, C-to-G, is 3 mismatch types) or Y # of mismatches, must be supplied as a integer vector of length 2. e.g. c(X, Y).
read_bqual  Exclude read if more than X percent of the bases have base qualities less than Y. Numeric vector of length 2. e.g. c(0.25, 20)
min_variant_reads  Required number of reads containing a variant for a site to be reported. Calculated per bam file, such that if 1 bam file has >= min_variant_reads, then the site will be reported.
min_allelic_freq  minimum allelic frequency required for a variant to be reported in ALT assay.
report_multiallelic  if TRUE, report sites with multiple variants passing filters. If FALSE, site will not be reported.
remove_overlaps  if TRUE, enable read pair overlap detection, which will count only 1 read in regions where read pairs overlap using the htslib algorithm. In brief for each overlapping base pair the base quality of the base with the lower quality is set to 0, which discards it from being counted.

Value

A RangedSummarizedExperiment object populated with multiple assays:

- ALT: Alternate base(s) found at each position
- nRef: # of reads supporting the reference base
- nAlt: # of reads supporting an alternate base
- nA: # of reads with A
- nT: # of reads with T
- nC: # of reads with C
- nG: # of reads with G
The `rowRanges()` contains the genomic interval for each site, along with:

- **REF**: The reference base
- **rpbz**: Mann-Whitney U test of Read Position Bias from bcftools, extreme negative or positive values indicate more bias.
- **vdb**: Variant Distance Bias for filtering splice-site artifacts from bcftools, lower values indicate more bias.
- **sor**: Strand Odds Ratio Score, strand bias estimated by the Symmetric Odds Ratio test, based on GATK code. Higher values indicate more bias.

The rownames will be populated with the format `site_[seqnames]_[position(1-based)]_[strand]`, with `strand` being encoded as 1 = +, 2 = -, and 3 = *.

**See Also**

Other pileup: `pileup_cells()`

**Examples**

```r
library(SummarizedExperiment)
bamfn <- raer_example("SRR5564269_Aligned.sortedByCoord.out.md.bam")
bam2fn <- raer_example("SRR5564277_Aligned.sortedByCoord.out.md.bam")
fafn <- raer_example("human.fasta")

rse <- pileup_sites(bamfn, fafn)
fp <- FilterParam(only_keep_variants = TRUE, min_depth = 55)
pileup_sites(bamfn, fafn, param = fp)

# using multiple bam files
bams <- rep(c(bamfn, bam2fn), each = 3)
sample_ids <- paste0(rep(c("KO", "WT"), each = 3), 1:3)
names(bams) <- sample_ids
fp <- FilterParam(only_keep_variants = TRUE)
rse <- pileup_sites(bams, fafn, param = fp)
rse

rse$condition <- substr(rse$sample, 1, 2)
assays(rse)

colData(rse)
rowRanges(rse)

# specifying regions to query using GRanges object
sites <- rowRanges(rse)
rse <- pileup_sites(bams, fafn, sites = sites)
```
raer

raer <- pileup_sites(bams, fafn, chroms = c("SPCS3", "DHFR"))
raer

raer <- pileup_sites(bams, fafn, region = "DHFR:100-101")
raer

raer: RNA editing tools in R

Description

Toolkit for identification and statistical testing of RNA editing signals from within R. Provides support for identifying sites from bulk-RNA and single cell RNA-seq datasets, and general methods for extraction of allelic read counts from alignment files. Facilitates annotation and exploratory analysis of editing signals using Bioconductor packages and resources.

Author(s)

Maintainer: Kent Riemondy <kent.riemondy@gmail.com> (ORCID)
Authors:

- Kristen Wells-Wrasman <kristen.wells-wrasman@cuanschutz.edu> (ORCID)

Other contributors:

- Ryan Sheridan <ryan.sheridan@cuanschutz.edu> (ORCID) [contributor]
- Jay Hesselberth <jay.hesselberth@gmail.com> (ORCID) [contributor]
- RNA Bioscience Initiative [copyright holder, funder]

See Also

Useful links:

- https://rnabioco.github.io/raer
- https://github.com/rnabioco/raer
- Report bugs at https://github.com/rnabioco/raer/issues
raer_example

*Provide working directory for raer example files.*

**Description**

Provide working directory for raer example files.

**Usage**

```r
raer_example(path)
```

**Arguments**

- `path`  
  path to file

**Value**

Character vector will path to an internal package file.

**Examples**

```r
raer_example("human.fasta")
```

---

read_sparray

*Read sparseMatrix produced by pileup_cells()*

**Description**

Read in tables produced by `pileup_cells()` which are an extension of the matrixMarket sparse matrix format to store values for more than 1 matrix.

The .mtx.gz files are formatted with columns:

1. row index (0 based)
2. column index (0 based)
3. values for sparseMatrix #1 (nRef)
4. values for sparseMatrix #2 (nAlt)

**Usage**

```r
read_sparray(mtx_fn, sites_fn, bc_fn, site_format = c("coordinate", "index"))
```
Arguments

- `mtx_fn` .mtx.gz file path
- `sites_fn` sites.txt.gz file path
- `bc_fn` bcs.txt.gz file path
- `site_format` one of `coordinate` or `index`, `coordinate` will populate a `SingleCellExperiment` with rowRanges and rownames corresponding to genomic intervals, whereas `index` will only add row indices to the rownames.

Value

a `SingleCellExperiment` object populated with `nRef` and `nAlt` assays.

Examples

```r
library(Rsamtools)
library(GenomicRanges)
bam_fn <- raer_example("5k_neuron_mouse_possort.bam")

gr <- GRanges(c("2:579:-", "2:625:-", "2:645:-", "2:589:-", "2:601:-"))
gr$REF <- c(rep("A", 4), "T")
gr$ALT <- c(rep("G", 4), "C")

cbs <- unique(scanBam(bam_fn, param = ScanBamParam(tag = "CB"))[[1]]$tag$CB)
cbs <- na.omit(cbs)

outdir <- tempdir()
bai <- indexBam(bam_fn)

fp <- FilterParam(library_type = "fr-second-strand")
mtx_fns <- pileup_cells(bam_fn, gr, cbs, outdir, return_sce = FALSE)
sce <- read_sparray(mtx_fns[1], mtx_fns[2], mtx_fns[3])
sce

unlink(bai)
```
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