

Package ‘LinTInd’

September 20, 2024

Title Lineage tracing by indels

Version 1.9.0

Description When we combine gene-editing technology and sequencing technology, we need to reconstruct a lineage tree from alleles generated and calculate the similarity between each pair of groups.

FindIndel() and IndelForm() function will help you align each read to reference sequence and generate scar form strings respectively.

IndelIdentifies() function will help you to define a scar form for each cell or read.

IndelPlot() function will help you to visualize the distribution of deletion and insertion.

TagProcess() function will help you to extract indels for each cell or read.

TagDist() function will help you to calculate the similarity between each pair of groups across the indwells they contain.

BuildTree() function will help you to reconstruct a tree.

PlotTree() function will help you to visualize the tree.

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| | |
|----------------|-----------------------|
| align_to_range | <i>align_to_range</i> |
|----------------|-----------------------|

Description

Title

Usage

align_to_range(p, s, cut)

Arguments

| | |
|-----|--|
| p | A base sequence in character format |
| s | A base sequence in character format |
| cut | The distance between the starting sites of two fragments |

Value

A list include two IRanges instances (deletion and insertion)

Examples

```
align_to_range(p="AAGG---AAATTCGGGAATAAGGAATTT",s="AAGGCCCAAATTT-CGGAATAAGGAATTT",cut=0)
```

 BuildTree

BuildTree

Description

Generate an array generant tree of a data.tree data structure and save it

Usage

```
BuildTree(tag)
```

Arguments

tag List generate from TagProcess, for more see [TagProcess](#)

Details

Title

Value

list with two elements, a data.tree data structure and a dataframe of array information for each cell barcode

Examples

```
data("example_data",package = "LinTInd")
treeinfo<-BuildTree(tag)
```

 cellsinfo

example of cellsinfo

Description

This an example of cellsinfo

Usage

```
data("example_data")
```

Format

cellsinfo

| | |
|----------|----------------------------|
| celltype | <i>example of celltype</i> |
|----------|----------------------------|

Description

This an example of celltype

Usage

```
data("example_data")
```

Format

celltype

| | |
|------------------|-------------------------|
| change_form_stat | <i>change_form_stat</i> |
|------------------|-------------------------|

Description

Title

Usage

```
change_form_stat(indel)
```

Arguments

| | |
|-------|---|
| indel | List include two IRanges instances, contain start and end site of deletions and inserstions |
|-------|---|

Value

A scar form string

Examples

```
data("example_data",package = "LinTInd")  
change_form_stat(cellsinfo$indel[[1]])
```

| | |
|---------|---------------------------|
| cutsite | <i>example of cutsite</i> |
|---------|---------------------------|

Description

This an example of cutsite

Usage

```
data("example_data")
```

Format

cutsite

| | |
|------|------------------------|
| data | <i>example of data</i> |
|------|------------------------|

Description

This an example of data

Usage

```
data("example_data")
```

Format

data

| | |
|--------------|---------------------|
| example_data | <i>example_data</i> |
|--------------|---------------------|

Description

A set of example data

Usage

```
data("example_data")
```

Format

The format is: chr "example_data"

Examples

```
data("example_data", package = "LinTInd")
```

FindIndel

FindIndel

Description

This function can identify indels for each read in input data, and create IRanges instances for deletion and insertion.

Usage

```
FindIndel(
  data,
  scarfull,
  scar,
  align_score = NULL,
  type = NULL,
  indel.coverage = NULL,
  cln
)
```

Arguments

| | |
|----------------|--|
| data | data frame, include cell barcode, UMI and reads. |
| scarfull | DNAStrng of reference sequence |
| scar | The cutsite data frame |
| align_score | The minimum alignment score that matched sequence should get, default in this parameter is the score that the reads which all of the target set were cutted got |
| type | Group name for this data ("None" in default) |
| indel.coverage | Choose indels selected scope: "Accurate" (default) means only the indels happened in target site will be identified; "All" means each indel will be detected even they locate on the anchors |
| cln | The number of threads |

Details

Title

Value

list include IRanges instances (deletion and insertion), a data frame of reads' informations, reference sequenc, dataframe of cut sites

Examples

```
data("example_data", package = "LinTInd")
scarinfo<-FindIndel(data=data, scarfull=ref, scar=cutsite, indel.coverage="All", type="test", cln=1)
```

IndelForm
*IndelForm***Description**

Generate scar form strings from scarinfo list for each reads

Usage

```
IndelForm(scarinfo, cln)
```

Arguments

| | |
|----------|--|
| scarinfo | List generate from FindIndel, for more see FindIndel |
| cln | The number of threads |

Details

Title

Value

A new list of scarinfo, the scarform of each reads will add in the data frame of reads' informations

Examples

```
data("example_data", package = "LinTInd")
IndelForm(scarinfo, cln=1)
```

IndelIdents
*IndelIdents***Description**

Function to define a scarform for each cell(single cell) or each reads(bulk seq, generate 'cell barcode' for each reads)

Usage

```
IndelIdents(scarinfo, method.use = NULL, cln)
```

Arguments

| | |
|------------|---|
| scarinfo | List generate from IndelForm, for more see IndelForm |
| method.use | Select how to determine a scar form string for each cell: "reads.num" (default):find the scar with the most reads in the cell; "umi.num":find the scar with the most UMIs in the cell; "consensus":find the consistent sequences in each cell, and then generate scar form strings from the new reads |
| cln | The number of threads |

Details

Title

Value

The list generate from FindIndel, but in 'Scar' element a new column contain scar form strings

Examples

```
data("example_data",package = "LinTInd")
IndelIdents(scarinfo,method.use="umi.num",cIn=1)
```

IndelPlot

IndelPlot

Description

Return 2 line charts, show the probability of insertion and deletion at each site

Usage

```
IndelPlot(cellsinfo)
```

Arguments

cellsinfo List generate from IndelIdents, for more see [IndelIdents](#)

Details

Title

Value

2 line charts

Examples

```
data("example_data",package = "LinTInd")
IndelPlot(cellsinfo = cellsinfo)
```

 PlotTree

PlotTree

Description

Function to visualise the array generate tree

Usage

```
PlotTree(treeinfo, data.extract = NULL, annotation = NULL, prefix = NULL)
```

Arguments

| | |
|--------------|---|
| treeinfo | List generate from BuildTree, for more see BuildTree |
| data.extract | (optional) If "FALSE" (default), will not return the indel's information, if it's "TRUE", the opposite |
| annotation | (optional) If "TRUE" (default), the annotation of each cell barcodes have to be provided before, and a heatmap of cells' distribution for each array will be return |
| prefix | (optional) Indels' prefix |

Details

Title

Value

A list include a ggplot object, a dataframe show the distribution of each array contained in each group of cells (optional), and a dataframe to create the ggplot object

Examples

```
data("example_data", package = "LinTInd")
plotinfo<-PlotTree(treeinfo = treeinfo,data.extract = "TRUE",annotation = "TRUE")
plotinfo<-PlotTree(treeinfo = treeinfo,data.extract = "TRUE",annotation = "FALSE")
```

 ReadCutsite

ReadCutsite

Description

Function to create a reference dataframe include each position and its' group

Usage

```
ReadCutsite(segref, reftype=NULL)
```

Arguments

| | |
|---------|---|
| segref | The cutsite file |
| reftype | Choose the reference type you want, if reftype="Accurate" (default), there will only the target sites be generated; if reftype="All", each site will be generated |

Details

Title

Value

reference dataframe

Examples

```
data("example_data", package = "LinTInd")
ReadCutsite(cutsite)
ReadCutsite(cutsite, reftype="All")
```

ReadFasta

ReadFasta

Description

Function to read fasta file to DNAStrng object

Usage

```
ReadFasta(filename)
```

Arguments

| | |
|----------|---------------------------|
| filename | The input fasta file name |
|----------|---------------------------|

Details

Title

Value

A DNAStrng object

Examples

```
fafile=paste0(system.file("extdata", package = 'LinTInd'), "/V3.fasta")
ReadFasta(fafile)
```

| | |
|-----|-----------------------|
| ref | <i>example of ref</i> |
|-----|-----------------------|

Description

This an example of ref

Usage

```
data("example_data")
```

Format

ref

| | |
|----------|----------------------------|
| scarform | <i>example of scarform</i> |
|----------|----------------------------|

Description

This an example of scarform

Usage

```
data("example_data")
```

Format

scarform

| | |
|----------|----------------------------|
| scarinfo | <i>example of scarinfo</i> |
|----------|----------------------------|

Description

This an example of scarinfo

Usage

```
data("example_data")
```

Format

scarinfo

| | |
|---------|---------------------------|
| scarref | <i>example of scarref</i> |
|---------|---------------------------|

Description

This an example of scarref

Usage

```
data("example_data")
```

Format

scarref

| | |
|-------------|-------------------------------|
| scarref_all | <i>example of scarref_all</i> |
|-------------|-------------------------------|

Description

This an example of scarref_all

Usage

```
data("example_data")
```

Format

scarref_all

| | |
|-----|-----------------------|
| tag | <i>example of tag</i> |
|-----|-----------------------|

Description

This an example of tag

Usage

```
data("example_data")
```

Format

tag

| | |
|---------|----------------|
| TagDist | <i>TagDist</i> |
|---------|----------------|

Description

If the cell barcode and the annotation of each cell are provided, this function can calculate the relationship between each cell type in three ways

Usage

```
TagDist(tag, method = NULL)
```

Arguments

- | | |
|--------|--|
| tag | List generated from TagProcess, for more see TagProcess |
| method | Denote which method to use: <ul style="list-style-type: none">• "Jaccard"(default): calculate the weighted jaccard similarity of indels between each pair of groups;• "P": right-tailed test, compare the Indels intersection level with the hypothetical result generated from random editing, and the former is expected to be significantly higher than the latter;• "spearman": Spearman correlation of indels between each pair of groups |

Details

Title

Value

2 figures are saved to show the distribution of INDEL and the relationship between groups respectively, the matrix of the relationship between groups is returned

Examples

```
tag_dist=TagDist(tag,method = "spearman")
tag_dist=TagDist(tag)
tag_dist=TagDist(tag,method = "Jaccard")
tag_dist=TagDist(tag,method = "P")
tag_dist=TagDist(tag,method = "spearman")
```

| | |
|------------|-------------------|
| TagProcess | <i>TagProcess</i> |
|------------|-------------------|

Description

Split each indel from scar form strings and map indel information to cell barcodes

Usage

```
TagProcess(data, Cells = NULL, prefix = NULL)
```

Arguments

| | |
|--------|--|
| data | List generate from IndelIdents, for more see IndelIdents |
| Cells | (optional) Dataframe of cells' annotation, with two columns: "Cell.BC" and "Cell.type" |
| prefix | (optional) Indels' prefix |

Details

Title

Value

List with two dataframes: Indels for each cell barcode and cells' annotation

Examples

```
data("example_data", package = "LinTInd")
TagProcess(cellsinfo$info, Cells=celltype)
```

| | |
|----------|----------------------------|
| treeinfo | <i>example of treeinfo</i> |
|----------|----------------------------|

Description

This an example of treeinfo

Usage

```
data("example_data")
```

Format

treeinfo

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