Package 'FMIndex'

September 7, 2023

Title A package for computing FM index
Version 0.99.0
Description The package computes the FM index of a DNA string contained in a fasta file. The user can specify the path where he wants to save the FM index data structures. The results will be also printed on the console.
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FMIndex-package FMIndex - Compute the FM index for a DNA sequence contained in a fasta file

Description

Package: FMIndex
Type: Package
Date: 2023-06-20
License: GPL (>=2)

Details

The FMIndex package implements all the functions necessary to compute the data structures of the FM index. The package has one main function which computes all the FM index data structures and saves them into different text files (the path is specified by the user):

- F_col.txt for the F column (cont the occurrences of the unique characters)
- L_col.txt for the L column (BWT)
- suffix_array.txt for the suffix array
- tally_table.txt for the tally table

The functions that compute the FM index data structures are available: they all take a DNAString as input and the return values are printed on the console.

NOTE: to make the strings compatible with DNAString objects the termination character used to build the BWT is "."

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FMindex	Writes the FM index	
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Description

This function writes the FM index of the DNA sequence contained in the FASTA file given in input

Usage

```
FMindex(inputData, usrpath, tallywidth = 1, includeEndChar = TRUE)
```

Arguments

inputData	Can be a DNAString or a FASTA file containing one single DNA sequence, files containing multiple sequences are not allowed.
usrpath	The directory path where the user wants to save the files: it must end with '/' (you can leave it empty if you want to save the files in your working directory).
tallywidth	How you want to compress the tally table. It states how often (in terms of rows) you save one checkpoint. Default value=1
includeEndChar	True if you want to include the "." in the F column, false if you want to exclude it. Default value:TRUE

Details

This function may raise an error if:

- The FASTA file provided contains more than one sequence
- The FASTA file provided contains no sequences

Value

A list containint all the data structures necessary for the FM index

Examples

```
FMindex(system.file("extdata/examples/seq.fasta",package="FMIndex"),"")
FMindex(system.file(
"extdata/examples/seq.fasta",package="FMIndex"),"",tallywidth=1)
```

4 getFcolumn

getBWTcharacter	Helper function: compute the i-th character of the BWT

Description

This is a helper function that performs a specific task for the main function getLcolumn. It computes the i-th character of the BWT of a given sequence which corresponds to the i-th character in the L column.

Usage

```
getBWTcharacter(index, sequence, suffixarray)
```

Arguments

index The position which you want to find the character

sequence The string you want to know the i-th L column character

suffixarray An integer vector containing the suffix array of the input string

Value

A character containing the i-th character of the input sequence

See Also

getLcolumn

Description

This function gets a DNAString and gives the F column of the FM index

Usage

```
getFcolumn(seq, includeEndChar = TRUE)
```

Arguments

seq the DNAString to be analysed

includeEndChar True if you want to include the ".", false if you want to exclude it. Default

value:TRUE

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Details

This function may raise a warning if

• The provided sequence is empty

Value

An integer vector with all the frequences in the dna sequence given in input

Examples

```
getFcolumn(Biostrings::DNAString("ACCT"))
```

getLcolumn

Compute the L column of a sequence

Description

This function gets a character and gives the F column of the FM index

Usage

```
getLcolumn(sequence)
```

Arguments

sequence

the DNAString to be analysed

Details

• The provided sequence is empty

Value

A list containing the L column (BWT) and the suffix array

Examples

```
getLcolumn(Biostrings::DNAString("ACGT"))
getLcolumn(Biostrings::DNAString("AACCGT"))
```

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 ${\tt getSA}$

Compute the suffix array of a string

Description

This function computes the suffix array of a given string

Usage

```
getSA(sequence)
```

Arguments

sequence

a DNAString representing the string sequence

Details

This function may raise a warning if

• The provided sequence is empty

Value

An integer array containing the suffix array

Examples

```
getSA(Biostrings::DNAString("ACGT"))
getSA(Biostrings::DNAString("AACCGT"))
```

getSuffix

Helper function: get the suffix

Description

This is a helper function that performs a specific task for the main function getSA.It gets the suffix of the string starting from a position given in input.

Usage

```
getSuffix(index, sequence)
```

Arguments

index An integer representing the starting position of the string

sequence The sequence you want to find the suffix

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Value

The substring starting at the given position

See Also

getSA

getTallyTable

Create the Tally Table

Description

Creates the Tally Table of a given BWT of a string

Usage

```
getTallyTable(1.col, rowwidth = 1)
```

Arguments

1. col A DNAString containing the L column of the FM index

rowwidth Step (in term of numer of rows) for saving checkpoints of the tally table. The

default value is 1 which corresponds to save all the rows of the tally table

Details

This function may raise an error if

- The argument rowwidth is higher than one
- The argument rowwidth is lower than one
- The input sequence (argument l.col) doesn't contain the termination character "."
- The input sequence (argument l.col) contains more than one termination character "."

This function may raise a warning if

• The input sequence contains only the termination character

Value

The tally table of the input sequence

Examples

```
getTallyTable(Biostrings::DNAString("T.AACCG"))
getTallyTable(Biostrings::DNAString("T.AACCG"),2)
```

8 getTallyTableLine

getTallyTableLine	Helper function: create the i-th row of the Tally Table	

Description

This is a helper function that performs a specific task for the main function getTallyTable. It creates the i-th row of the Tally Table

Usage

```
getTallyTableLine(index, 1.col, alphabet)
```

Arguments

index An integer representing the index of the i-th row you want to build

1.col A DNAString object containing the L column of the FM index

alphabet A character vector containing the alphabet to consider in counting the elements

Value

the i-th row of the Tally Table

See Also

getTallyTable

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