# Package 'FMIndex'

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Title A package for computing FM index
<b>Version</b> 0.99.0
<b>Description</b> 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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Suggests knitr, rmarkdown, BiocStyle, testthat
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FMIndex-package

FMIndex - Computer the FM index for a DNA sequence contained in a fasta file The FMIndex package implements all the functions necessary to compute the data structures of the FM index

## Description

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

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#### References

http://some.url/here/maybe/
Resta et al.

**FMindex** 

Writes the FM index

## Description

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

## Usage

FMindex(fastafile, usrpath, tallywidth = 1, includeTerminationCharacter = TRUE)

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#### **Arguments**

fastafile The .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. Default value=1

includeTerminationCharacter

default value: true

## Value

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

## Examples

```
FMindex(system.file("extdata/examples/seq5.fasta",package="FMIndex"),"")
FMindex(system.file(
"extdata/examples/seq5.fasta",package="FMIndex"),"",tallywidth=1)
```

getBWTcharacter

Compute the i-th character of the BWT

#### **Description**

This function 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

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getFcolumn

Compute the F column

#### **Description**

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

## Usage

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

#### **Arguments**

seq the DNAString to be analysed

include Termination Character

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

value:TRUE

#### 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 string to be analysed

### Value

A DNAString containing the L column of the FM index

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#### **Examples**

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

getSuffix

Get the suffix

## **Description**

This function 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

#### Value

The substring starting at the given position

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 for saving rows of the tally table. The default value is 1 which corresponds

to save all the rows of the tally table

#### Value

The tally table of the input sequence

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## **Examples**

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

getTallyTableLine

Create the i-th row of the Tally Table

## Description

This function 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

## **Index**

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