Package 'GenomicTools.fileHandler'

October 12, 2022

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GenomicTools.fileHandler-package

R Package To Handle Files From Genomic Data Genomic-Tools.fileHandler is a loose collection of I/O Functions Needed in Genomic Data Analysis

Description

Package: GenomicTools.fileHandler

Type: Package Version: 0.1.5.9 Date: 2020-03-05 example.bed 3

License: GPL LazyLoad: yes

Author(s)

Daniel Fischer

Maintainer: Daniel Fischer <daniel.fischer@luke.fi>

example.bed

Example Gene Annotation in Bed-Format

Description

This file contains some example lines to represent a typical bed file that can be used to try the corresponding functions.

Format

A file with three column Chr, Start and End.

Details

The file is locate din the /extdata folder of the package and is accessible after installation via system.file("extdata", "example.bed", package="GenomicTools.fileHandler")

Author(s)

Daniel Fischer

example.fasta

Example Sequencing Reads in fasta-Format

Description

This file contains some example reads to represent a typical fasta file that can be used to try the corresponding functions.

Details

The file is locate din the /extdata folder of the package and is accessible after installation via system.file("extdata", "example.fasta", package="GenomicTools.fileHandler")

Author(s)

4 example.gff

example.fastq

Example Sequencing Reads in fastq-Format

Description

This file contains some example reads to represent a typical fastq file that can be used to try the corresponding functions.

Details

The file is locate din the /extdata folder of the package and is accessible after installation via system.file("extdata", "example.fastq", package="GenomicTools.fileHandler")

Author(s)

Daniel Fischer

example.gff

Example Gene Annotation in gff-Format

Description

This file contains some example gene annotations to represent a typical gff file that can be used to try the corresponding functions.

Details

The file is locate din the /extdata folder of the package and is accessible after installation via system.file("extdata", "example.gff", package="GenomicTools.fileHandler")

Author(s)

example.gtf 5

example.gtf

Example Gene Annotation in gtf-Format

Description

This file contains some example gene annotations to represent a typical gtf file that can be used to try the corresponding functions.

Details

The file is locate din the /extdata folder of the package and is accessible after installation via system.file("extdata", "example.gtf", package="GenomicTools.fileHandler")

Author(s)

Daniel Fischer

example.ped

Example Variant data in ped/map-Format

Description

This file contains some example variants to represent a typical ped/map file pair that can be used to try the corresponding functions.

Details

The file is locate din the /extdata folder of the package and is accessible after installation via system.file("extdata", "example.ped", package="GenomicTools.fileHandler")

Author(s)

6 example2.gtf.gz

example.vcf

Example Variant data in vcf-Format

Description

This file contains some example variants to represent a typical vcf file that can be used to try the corresponding functions.

Details

The file is locate din the /extdata folder of the package and is accessible after installation via system.file("extdata", "example.vcf", package="GenomicTools.fileHandler")

Author(s)

Daniel Fischer

example2.gtf.gz

Example Gene Annotation in zipped gtf-Format

Description

This file contains some example gene annotations to represent a typical zipped gtf file that can be used to try the corresponding functions.

Details

The file is locate din the /extdata folder of the package and is accessible after installation via system.file("extdata", "example2.gtf.gz", package="GenomicTools.fileHandler")

Author(s)

exportBed 7

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Exporting a Bed File.

Description

This function exports a standard bed file.

Usage

```
exportBed(x, file = NULL, header = FALSE)
```

Arguments

x data.frame

file Character, specifies filename/path header Logical, shall a header be written

Details

This function exports a data.frame to a standard bed file. If no file name is given, the variable name will be used instead.

Value

A bed file

Author(s)

Daniel Fischer

Examples

8 exportFA

exportFA

Exporting a Fasta File.

Description

This function exports a standard fasta file.

Usage

```
exportFA(fa, file = NULL)
```

Arguments

fa fasta object

file Character, specifies filename/path

Details

This function exports a fasta object to a standard fasta file. If no file name is given, the variable name will be used instead.

Value

A fasta file

Author(s)

Daniel Fischer

Examples

```
# Define here the location on HDD for the example file
  fpath <- system.file("extdata","example.fasta", package="GenomicTools.fileHandler")
# Import the example fasta file
  fastaFile <- importFA(file=fpath)
  newFasta <- fastaFile[1:5]

myfile <- file.path(tempdir(), "myLocs.fa")
  exportFA(newFasta, file=myfile)</pre>
```

importBed 9

Description

This function imports a standard bed file

Usage

```
importBed(file, header = FALSE, sep = "\t")
```

Arguments

file Specifies the filename/path header Logical, is a header present

sep Column separator

Details

This function imports a standard bed-file into a data.frame. It is basically a convenience wrapper around read.table. However, if no header lines is given, this function automatically assigns the column names, as they are given in the bed-specification on the Ensembl page here: https://www.ensembl.org/info/website/upload/bed.html

Value

```
A data.frame
```

Author(s)

Daniel Fischer

See Also

```
[exportBed], [read.table]
```

Examples

```
# Define here the location on HDD for the example file
  fpath <- system.file("extdata","example.bed", package="GenomicTools.fileHandler")
# Import the example bed file
  bedFile <- importBed(file=fpath)</pre>
```

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importBlastTab

Import a Tab Delimited Blast Output File

Description

This function imports a tab delimited blast output.

Usage

```
importBlastTab(file)
```

Arguments

file

Filename

Details

This function imports a tab delimited blast output file, currently the same as read. table

Value

A data.frame

Author(s)

Daniel Fischer

importFA

Importing a Fasta File.

Description

This function imports a standard fasta file

Usage

```
importFA(file)
```

Arguments

file

Specifies the filename/path

importFeatureCounts 11

Details

This function imports a standard fasta file. Hereby, it does not matter if the identifier and sequence are alternating or not, as the rows starting with '>' are used as identifier.

The example file was downloaded from here and was then further truncated respective transformed to fasta format:

ftp://ftp.1000genomes.ebi.ac.uk/vol1/ftp/phase3/data/HG00096/sequence_read/

Value

An object of class fa containing the sequences. The names correspond to the sequence names given in the fasta file.

Author(s)

Daniel Fischer

See Also

print.fa, summary.fa

Examples

```
# Define here the location on HDD for the example file
  fpath <- system.file("extdata","example.fasta", package="GenomicTools.fileHandler")
# Import the example fasta file
  fastaFile <- importFA(file=fpath)</pre>
```

importFeatureCounts

Import from FeatureCounts

Description

This functions imports the output from FeatureCounts

Usage

```
importFeatureCounts(file, skip = 0, headerLine = 2)
```

Arguments

file Character, file name

skip Number of lines to skip from txt file

headerLine Linenumber that contains the header information

importFQ

Details

FeatureCounts produces two files, the txt that contain the expression values and then the summary that containts all the information about the mapping statistics. This function imports both and stores them in a corresponding list.

Value

A list with expValues, geneInfo and summary

Author(s)

Daniel Fischer

Examples

```
# Define here the location on HDD for the example file
  fpath <- system.file("extdata","featureCountsExample.txt", package="GenomicTools.fileHandler")
# Import the example featureCounts file
  fcFile <- importFeatureCounts(file=fpath)</pre>
```

importFQ

Importing a Fastq File.

Description

This function imports a standard fastq file

Usage

```
importFQ(file)
```

Arguments

file

Specifies the filename/path

Details

This function imports a standard fastq file that consists out of blocks of four lines per entry

Value

An object of class fq containing the sequences and the quality measure. The names correspond to the sequence names given in the fasta file.

Author(s)

importGFF 13

See Also

```
print.fq, summary.fq
```

Examples

```
# Define here the location on HDD for the example file
  fpath <- system.file("extdata","example.fastq", package="GenomicTools.fileHandler")
# Import the example fastq file
  fastqFile <- importFQ(file=fpath)</pre>
```

importGFF

importGFF

Description

Import a GFF file

Usage

```
importGFF(file, skip = "auto", nrow = -1, use.data.table = TRUE,
  level = "gene", features = NULL, num.features = c("FPKM", "TPM"),
  print.features = FALSE, merge.feature = NULL, merge.all = TRUE,
  class.names = NULL, verbose = TRUE)
```

Arguments

file file or folder skip numeric, lines to skip numeric, lines to read nrow use.data.table logical level Character, read level, default: "gene" features features to import num.features names of the numeric features print.features Logical, print available features Character, merge multiple samples to dataset merge.feature merge.all Logical, shall all samples be merged together Definition of class name sin V9 class.names verbose Logical, verbose function output

Details

This function imports a standard gff file.

importGFF3

Value

A gff object

Author(s)

Daniel Fischer

Examples

```
# Define here the location on HDD for the example file
  fpath <- system.file("extdata","example.gff", package="GenomicTools.fileHandler")
# Import the example gff file
  importGFF(fpath)</pre>
```

importGFF3

importGFF3

Description

Import a GFF3 file

Usage

```
importGFF3(gff, chromosomes)
```

Arguments

gff file or folder

chromosomes The chromosome to import

Details

This function imports a standard gff3 file.

Value

A gff object

Author(s)

importGTF 15

importGTF	Import a GTF File	
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Description

This function imports a gtf file.

Usage

```
importGTF(file, skip = "auto", nrow = -1, use.data.table = TRUE,
  level = "gene", features = NULL, num.features = c("FPKM", "TPM"),
  print.features = FALSE, merge.feature = NULL, merge.all = TRUE,
  class.names = NULL, verbose = TRUE)
```

Arguments

file file or folder

skip numeric, lines to skip nrow numeric, lines to read

use.data.table logical

level Character, read level, default: "gene"

features features to import

num.features names of the numeric features print.features Logical, print available features

merge.feature Character, merge multiple samples to dataset

merge.all Logial, shall all samples be merged

class.names Vector with class names

verbose Logical, verbose function output

Details

This function imports a gtf file. The features names to be imported are defined in features, several features are then provided as vector. A list of available feature can be printed, by setting print.features=TRUE.

The skip option allows to skip a given number of rows, the default is, however, auto. In that case, all rows that start with the # symbol are skipped.

In case a set of expression values given in gtf format should be imported and to be merged into a single data table, the feature that should be used for merging can be provided to the merge. feature option. In that case the function expects a folder in file and it will import all gtfs located in that folder and merges them according to the merge. feature option. With the option class.names a vector of prefixes for the merged features can be provided. If this is kept empty, then the filenames of the gtf will be used instead (without gtf extension).

By default the function imprts all features in column 9 as string character. However, for common labels (FPKM and TPM) the class type is set automatically to numeric. Additional numerical feature names can be defined with the num. feature option.

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Value

A gtf object

Author(s)

Daniel Fischer

Examples

```
# Define here the location on HDD for the example file
  fpath <- system.file("extdata","example.gtf", package="GenomicTools.fileHandler")
# Same file, but this time as gzipped version
  fpath.gz <- system.file("extdata","example2.gtf.gz", package="GenomicTools.fileHandler")
# Import the example gtf file
  importGTF(fpath, level="transcript", features=c("gene_id","FPKM"))
## Not run:
# For the current you need to have zcat installed (should be standard on a Linux system)
  importGTF(fpath.gz, level="transcript", features=c("gene_id","FPKM"))
## End(Not run)</pre>
```

importPED

importPED

Description

Import a PED/MAP file pair

Usage

```
importPED(file, n, snps = NULL, which, split = "\t| +", sep = ".",
na.strings = "0", lex.order = FALSE, verbose = TRUE)
```

Arguments

file

n	Number of samples to read
snps	map filename
which	Names of SNPS to import
split	Columns separator in ped file
sep	Character that separates Alleles
na.strings	Definition for missing values
lex.order	Logical, lexicographical order
verbose	Logical, verbose output

ped filename

importSTARLog 17

Details

This function is to a large extend taken from snpStat::read.pedmap, but here is internally the data.table::fread function used that resulted in much faster file processing.

To import the data, the ped file can be provided to the file option and the map file to the snps option. If no option is given to snps and the file option is provided without any file extension, then the ped/map extension are automaticall added

Value

```
a pedmap object
```

Author(s)

Daniel Fischer

Examples

```
# Define here the location on HDD for the example file
  pedPath <- system.file("extdata","example.ped", package="GenomicTools.fileHandler")
  mapPath <- system.file("extdata","example.map", package="GenomicTools.fileHandler")
# Import the example ped/map files
  importPED(file=pedPath, snps=mapPath)</pre>
```

importSTARLog

importSTARLog

Description

Import the Log-File from STAR

Usage

```
importSTARLog(dir, recursive = TRUE, log = FALSE, finalLog = TRUE,
  verbose = TRUE)
```

Arguments

dir	The directory name
recursive	Logical, check for sub-directories
log	boolean, import also log file
finalLog	boolean, import also final_log file
verbose	Logical, talkactive function feedback

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Details

This function imports the Log file from STAR

Value

a data frame

Author(s)

Daniel Fischer

importVCF

importVCF

Description

Import a VCF function

Usage

```
importVCF(file, na.seq = "./.")
```

Arguments

file The file name

na.seq The missing value definition

Details

This function imports a VCF file.

In case the logic flag 'phased' is set to TRUE then the genotypes are expected to be in the format 010, otherwise they are expected to be like 0/1.

The example file was downloaded from here:

ftp://ftp.1000genomes.ebi.ac.uk/vol1/ftp/pilot_data/release/2010_07/exon/snps/

Value

A vcf object

Author(s)

importXML 19

Examples

```
# Define here the location on HDD for the example file
  fpath <- system.file("extdata","example.vcf", package="GenomicTools.fileHandler")
# Import the example vcf file
  importVCF(fpath)</pre>
```

importXML

importXML

Description

Import an Blast XML file

Usage

```
importXML(folder, seqNames = NULL, which = NULL, idTH = 0.8,
  verbose = TRUE)
```

Arguments

folder Character, folder path
seqNames Names of sequences
which Which sequences to import
idTH Use the threshold as cut-off
verbose Logical, verbose output

Details

This function imports XML files as provided as Blast output, it is mainly aimied to import the output of the hoardeR package

Value

An XML object

Author(s)

plotTotalReads

plotTotalReads

Description

Plot the total reads

Usage

plotTotalReads(STARLog)

Arguments

STARLog

A STARLog object

Details

This function plots the total reads from a STARlog object

Part of the diagnostic plot series for of the STARLog. The function accepts also a list of STARLogs and creates then comparative boxplots

Value

A plot

Author(s)

Daniel Fischer

 $\verb"plotUniquelyMappedReads"$

plot Uniquely Mapped Reads

Description

Plot the uniquely mapped reads

Usage

plotUniquelyMappedReads(STARLog)

Arguments

STARLog

A STARLog object

prereadGTF 21

Details

This function plots the percenage of uniquely reads from a STARlog object

Part of the diagnostic plot series for of the STARLog. The function accepts also a list of STARLogs and creates then comparative boxplots

Value

A plot

Author(s)

Daniel Fischer

prereadGTF

prereadGTF

Description

Preread a gtf file and prints features of it for importing it.

Usage

```
prereadGTF(file, nrow = 1000, skip = "auto")
```

Arguments

file Filename

nrow Number of rows to read skip Rows to skip from top

Details

This function reads in a gtf file and prints its features for the import step.

By default this function only imports the first 1000 rows, in case all rows should be imported set nrow=-1.

The number to skip in the beginning can be adjusted by the skip option. The default is here auto so that the function can identify the correct amount of header rows. Hence, this option should be changed only, if there is a good reason.

Value

A list of available features

Author(s)

22 print.fa

print.bed

Print a bed Object

Description

Prints a bed object.

Usage

```
## S3 method for class 'bed'
print(x, n = 6, ...)
```

Arguments

x Object of class bed.n Number of lines to print... Additional parameters

Details

The print function displays a bed object

Author(s)

Daniel Fischer

print.fa

Print a fa Object

Description

Prints a fa object.

Usage

```
## S3 method for class 'fa'
print(x, n = 2, seq.out = 50, ...)
```

Arguments

X	Object of class fa.
n	Number of sequences to display
seq.out	Length of the subsequence to display
	Additional parameters

print.featureCounts 23

Details

The print function displays a fa object

Author(s)

Daniel Fischer

print.featureCounts

Print a featureCounts Object

Description

Prints an featureCounts object.

Usage

```
## S3 method for class 'featureCounts' print(x, ...)
```

Arguments

x Object of class featureCounts.

... Additional parameters

Details

The print function displays a featureCounts object

Author(s)

Daniel Fischer

print.fq

Print a fq Object

Description

Prints a fq object.

Usage

```
## S3 method for class 'fq'
print(x, n = 2, seq.out = 50, print.qual = TRUE, ...)
```

24 print.gtf

Arguments

x Object of class fq.

n Number of sequences to display

seq.out Length of the subsequence to display

print.qual Logical, shall the quality measures also be printed

... Additional parameters

Details

The print function displays a fa object

Author(s)

Daniel Fischer

print.gtf

Print a gtf Object

Description

Prints a gtf object.

Usage

```
## S3 method for class 'gtf'
print(x, n = 6, ...)
```

Arguments

x Object of class gtf.

n Number of lines to print

... Additional parameters

Details

The print function displays a bed object

Author(s)

print.pedMap 25

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Print a pedMap Object

Description

Prints an pedMap object.

Usage

```
## S3 method for class 'pedMap'
print(x, n = 6, m = 6, ...)
```

Arguments

x Object of class pedMap.

Number of samples to displayNumber of columns to display

... Additional parameters

Details

The print function displays a pedMap object

Author(s)

Daniel Fischer

print.vcf

Print a vcf Object

Description

Prints an vcf object.

Usage

```
## S3 method for class 'vcf'
print(x, n = 6, m = 6, fullHeader = FALSE, ...)
```

Arguments

ject of class vcf.
)

n Number of samples to displaym Number of columns to display

fullHeader Logical, shall the whole header be printed

... Additional parameters

26 summary.fa

Details

The print function displays a vcf object

Author(s)

Daniel Fischer

summary.bed

Summary of a bed Object

Description

Summarizes a bed object.

Usage

```
## S3 method for class 'bed'
summary(object, ...)
```

Arguments

object Object of class bed.... Additional parameters

Details

The summary function displays an informative summary of a bed object

Author(s)

Daniel Fischer

summary.fa

Summary of a fa Object

Description

Summarizes a fa object.

Usage

```
## S3 method for class 'fa'
summary(object, ...)
```

summary.featureCounts 27

Arguments

```
object Object of class fa.... Additional parameters
```

Details

The summary function displays an informative summary of a fa object

Author(s)

Daniel Fischer

```
summary. feature Counts \ \textit{Summary of a feature Counts Object}
```

Description

Summarizes a featureCounts object.

Usage

```
## S3 method for class 'featureCounts'
summary(object, ...)
```

Arguments

object Object of class featureCounts.

... Additional parameters

Details

The summary function displays an informative summary of a featureCounts object

Author(s)

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summary.fq

Summary of a fq Object

Description

Summarizes a fq object.

Usage

```
## S3 method for class 'fq'
summary(object, ...)
```

Arguments

object Object of class fq.... Additional parameters

Details

The summary function displays an informative summary of a fq object

Author(s)

Daniel Fischer

summary.gtf

Summary of a gtf Object

Description

Summarizes a gtf object.

Usage

```
## S3 method for class 'gtf'
summary(object, ...)
```

Arguments

```
object Object of class gtf.... Additional parameters
```

Details

The summary function displays an informative summary of a gtf object

summary.STARLog 29

Author(s)

Daniel Fischer

summary.STARLog

Summary of a STARLog Object

Description

Summarizes a STARLog object.

Usage

```
## S3 method for class 'STARLog'
summary(object, ...)
```

Arguments

object Object of class STARLog.
... Additional parameters

Details

The summary function displays an informative summary of a STARLog object

Author(s)

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