Rsubread

October 25, 2011

align

Align next-gen sequencing reads to reference genome

Description

This is a R wrapper function for aligning reads. This function calls the underlying C function

Usage

align(index,readfile1,readfile2=NULL,output_file,nsubreads=10,TH1=3,TH2=3,nthrea

Arguments

index	character string giving the basename of index file. Index files should be located in the current directory.
readfile1	character string giving the name of file which includes sequencing reads. This will be the name of first file of paired-end data are provided. The read file should be in FASTQ or FASTA format.
readfile2	character string giving the name of second file when paired-end read data are provided. NULL by default.
output_file	character string giving the name of output file which includes the mapping results.
nsubreads	numeric value giving the number of subreads extracted from each read. 10 by default.
TH1	numeric value giving the consensus threshold for reporting a hit. This threshold will be applied to the first read if paired-end read data are provided. 3 by default.
TH2	numeric value giving the consensus threhold for the second read in a pair. 3 by default.
nthreads	numeric value giving the number of threads used for mapping. 1 by default.
indels	numeric value giving the number of insertions/deletions allowed during the mapping. 0 by default.
min_distance	numeric value giving the minimal distance between the pair of reads. 50 by default.
max_distance	numeric value giving the maximal distance between the pair of reads. 500 by default.

PE_orientation

character string giving the orientation of the two reads in a pair. "fr" by default, which means the first read is on the forward strand and the second read is on the reverse strand.

Details

This function implements a novel mapping strategy which uses a set of 16bp substrings (called subreads) extracted from each read to map them to the reference genome. Different from "seed-and-extend" mapping strategy, this new strategy does not have an extension step therefore it is a lot faster than the competing aligners. Due to the short length of the selected subsreads (16bp long), this strategy has a much high sensitivity than other aligners (seed length is usually around 30bp), i.e. it can align a lot more reads than competing aligners. Our evaluation results (using both simulation dataset and real dataset) showed that the accuracy of the new strategy is comparable to or slightly better than other aligners.

Two key parameters used by this new strategy are the number of subreads selected nsubreads and the consensus threshold for determining mapping locations TH1 (also TH2 for paired-end read data). We recommend using the default setting of these parameters to map reads of around 100bp long. However, users can choose to use more subreads when mapping longer reads. We recommend to set the value of consensus threshold to be 30 percent of the number of subreads used.

The C implementation of this strategy can be found at http://sourceforge.net/projects/subread/. This R function calls the corresponding C function to perform the alignment. Therefore, it has the mapping speed as the C program.

buildindex function should be called if the index has not been built for the reference genome. The index can be re-used once it has been built.

If paired-end read data is provided, file readfile1 will assumed to contain the first read from the read pair and readfile2 the second read.

Value

A file of SAM format which includes the mapping results.

Author(s)

Wei Shi and Yang Liao

References

Yang Liao and Wei Shi, "Subread: a superfast read aligner with high sensitivity and accuracy", In preparation.

```
library(Rsubread)
ref <- system.file("extdata", "reference.fa", package="Rsubread")
path <- system.file("extdata", package="Rsubread")
buildindex(basename=file.path(path, "reference_index"), reference=ref)
reads <- system.file("extdata", "reads.txt", package="Rsubread")
align(index=file.path(path, "reference_index"), readfile1=reads, output_file=file.path(path, "path")</pre>
```

atgcContent

Description

Calculate percentages of nucletodies A, T, G and C

Usage

atgcContent(filename, basewise=FALSE)

Arguments

filename	character string giving the name of input FASTQ/FASTA file
basewise	logical. If TRUE, nucleotide percentages will be calculated for each base position in the read across all the reads. By default, percentages are calculated for the entire dataset.

Details

Sequencing reads could contain letter "N" besides "A", "T", "G" and "C". Percentage of "N" in the read dataset is calcuated as well.

The basewise calculation is useful for examining the GC bias towards the base position in the read. By default, the percentages of nucleotides in the entire dataset will be reported.

Value

A named vector containing percentages for each nucleotide type if basewise is FALSE. Otherwise, a data matrix containing nucleotide percentages for each base position of the reads.

Author(s)

Zhiyin Dai and Wei Shi

```
library(Rsubread)
reads <- system.file("extdata","reads.txt",package="Rsubread")
# Fraction of A,T,G and C in the entire dataset
x <- atgcContent(filename=reads,basewise=FALSE)
# Fraction of A,T,G and C at each base location across all the reads
xb <- atgcContent(filename=reads,basewise=TRUE)</pre>
```

```
buildindex
```

Description

This is an R wrapper function for building index for a reference genome. This function calls the underlying C function.

Usage

```
buildindex (basename, reference, colorspace=FALSE, memory=3700)
```

Arguments

basename	character string giving the basename of created index files.
reference	charater string giving the name of the file containing all the refernece sequences.
colorspace	logical. If ${\tt TRUE},$ a color space index will be built. Otherwise, a base space index will be built.
memory	numeric value specifying the amount of memory to be requested in gigabytes

Details

A hash table will be built for the reference genome. Keys in the hash table are the 16bp sequences and hash values are their chromosomal locations. A 16bp sequence could have one or more than one chromosomal locations. They are all recorded in the hash table. Non-informative 16bp sequences, which are highly repetitive in the reference genome, are not included in the hash table.

After the index is built, reads can then be mapped to the reference genome by using align function.

It takes around 1 hour to build an index for human genome.

Value

Index files with basename provided in basename. These files are saved in the current directory.

Author(s)

Wei Shi and Yang Liao

```
library(Rsubread)
ref <- system.file("extdata", "reference.fa", package="Rsubread")
path <- system.file("extdata", package="Rsubread")
buildindex(basename=file.path(path, "reference_index"), reference=ref)</pre>
```

featureCounts Count the number of mapped reads for each feature

Description

Summarize read counts to features including genes and exons

Usage

```
featureCounts(SAMfiles,type="gene",species="mm",annot=NULL)
```

Arguments

SAMfiles	a character vector giving names of SAM format files.
type	a character string giving the feature type. Its value could be gene or exon.
species	a character string specifying the species. It can be mm or hg. Values of this argument determines which in-built annotation file will be used, if annot is NULL.
annot	a character string giving the name of the annotation file provided by users, which includes feature information such as chromosomal coordinates etc. This file will override the in-built annotation file chosen from using 'species' argument.

Details

This function takes as input a set of SAM format files and assigns reads to the features. Currently, only feature types including gene and exon are supported. gene is the aggregation of all the exons for each gene.

There are two in-built annotation files which are used by this function to summarize reads for genes or exons for mouse and human, respectively. These annotation files include the exon annotation information downloaded from NCBI Build 37.2, including Entrez gene identifier and chromosomal coordinates for each exon. The species argument specifies which annotation file should be used.

Users can provide their own annotation file for read summarization as well, by using the annot argument. In this case, the user provided annotation file will override the in-built annotation file. The annotation file provided by users should be a tab delimited file, and its first four columns should provide gene identifiers, chromosome names, chromosomal start locations and chromosomal end locations for each exon, respectively. Below is an example:

```
entrezid chromosome chr_start chr_stop
497097 chr1 3204563 3207049
497097 chr1 3411783 3411982
497097 chr1 3660633 3661579
100503874 chr1 3637390 3640590
100503874 chr1 3648928 3648985
100038431 chr1 3670236 3671869
```

• • •

Although this function is designed for summarizing reads from RNA-seq experiments, it can be used to summarize reads from other next-gen sequencing experiments as well, for example ChIP-seq or other DNA sequencing experiments. Simply by setting type to exon and providing an annotation, this function will yield numbers of mapped reads for each feature.

A data frame containing read counts for each feature.

Author(s)

Wei Shi

propmapped

Obtain the proportion of mapped reads

Description

Use mapping information stored in a SAM format file to count the number of mapped reads

Usage

```
propmapped(samfile)
```

Arguments

samfile character string giving the name of a SAM format file.

Details

This function uses the mapping information included in a SAM format file get the proportion mapped reads out of all the reads.

Value

Fraction of mapped reads is printed on the screen.

Author(s)

Wei Shi

```
library(Rsubread)
results <- system.file("extdata","alignResults.SAM",package="Rsubread")
propmapped(results)</pre>
```

qualityScores

Description

Extract quality scores and convert them to ASCII code

Usage

qualityScores(filename, offset=64, nreads=10000)

Arguments

filename	character string giving the name of input FASTQ file.
offset	numeric value giving the offset added to the original quality score, 64 by default.
nreads	numeric value giving the number of reads from which quality scores are ex- tracted

Details

Quality scores are given in the form of characters in datasets which contain sequencing reads. This function extracts the quality scores and then convert them to the ASCII codes which encode these characters. These ASCII codes are then subtracted by the offset to obtain the original quality scores.

If the total number of reads is n, then every n/nreads read will be used for quality score retrieval.

Value

A data matrix containing the quality scores with rows being reads and columns being base positions in the read.

Author(s)

Zhiyin Dai and Wei Shi

```
library(Rsubread)
reads <- system.file("extdata","reads.txt",package="Rsubread")
x <- qualityScores(filename=reads,nreads=1000)
boxplot(x)</pre>
```

sam2bed

Description

SAM to BED conversion

Usage

```
sam2bed(samfile,bedfile,readlen)
```

Arguments

samfile	character string giving the name of input file. Input format should be in SAM format.
bedfile	character string giving the name of output file. Output file is in BED format.
readlen	numeric value giving the length of reads included in the input file.

Details

SAM format is the de facto standard format of output from read aligner. This format not only includes the mapping coordinates of the reads but also includes other using information such as mapping quality, CIGAR information and so on. This function converts a SAM format file to a BED format file, which can then be displayed in a genome browser like UCSC genome browser, IGB, IGV etc.

Value

A BED format file.

Author(s)

Wei Shi

```
library(Rsubread)
results <- system.file("extdata", "alignResults.SAM", package="Rsubread")
sam2bed(samfile=results, bedfile="alignResults.bed", readlen=100)</pre>
```

Index

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