Package ‘ChIPsim’

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Description A general framework for the simulation of ChIP-seq data. Although currently focused on nucleosome positioning the package is designed to support different types of experiments.
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ChIPsim-package .................................................. 2
bindDens2readDens .............................................. 3
decodeQuality .................................................... 4
defaultControl ................................................... 5
defaultErrorProb ................................................ 6
defaultFunctions ............................................... 6
defaultGenerator .............................................. 7
extractQuality ................................................... 8
feat2dens .......................................................... 9
featureDensity .................................................. 10
FeatureGenerators ............................................. 11
makeFeatures ................................................... 12
placeFeatures .................................................. 13
pos2fastq ......................................................... 16
readError ......................................................... 17
Description

This package provides a framework for the simulation of ChIP-seq experiments. An implementation of a simulation for nucleosome positioning experiments is provided as part of the package. Simulations for other experiments can be implemented using the provided framework.

Details

Function simChIP is the main driver of the simulation. To simulate different types of experiments the functions passed to the functions argument of simChIP have to be replaced. See the vignettes for more detail.

Author(s)

Peter Humburg
Maintainer: Peter Humburg <Peter.Humburg@well.ox.ac.uk>

References

~~ Literature or other references for background information ~~

See Also

ShortRead and its dependencies are used to handle short read and genomic sequences.
Examples

## See the accompanying vignette 'Introduction to ChIPsim' for a detailed
## example of how to use this package for nucleosome positioning simulations.
## A guide for the writing of extensions that cover other types of
## experiments is provided in 'Extending ChIPsim'.

bindDens2readDens

### Convert a feature density into a read density

Description

Given a feature density this function produces two read densities, one for each strand.

Usage

`bindDens2readDens(bindDens, fragment, nfrag = 1e+05, bind = 147,
minLength = 150, maxLength = 180, ...)

Arguments

bindDens Numeric vector with the feature density for one chromosome.
fragment Function giving the fragment length distribution.
nfrag Number of fragments that should be simulated to generate the read distribution.
bind Length of binding site.
minLength Minimum fragment length.
maxLength Maximum fragment length.
...
Further arguments to `fragment`.

Value

A two column matrix. The first column contains the read density for the forward strand, the second column the read density for the reverse strand.

Author(s)

Peter Humburg

See Also

`feat2dens`, `sampleReads`

Examples

```r
set.seed(1)
## generate a (relatively short) sequence of nucleosome features
features <- placeFeatures(start=200, length=1e5)

## calculate feature density
featureDens <- feat2dens(features, length=1e5)

## convert to read density
readDens <- bindDens2readDens(featureDens, fragDens, meanLength=160)
```
## decodeQuality

### Conversion between numerical and ASCII representation of read qualities

#### Description

These functions convert an ASCII encoded sequence of read qualities into a numeric vector of error probabilities and vice versa.

#### Usage

```r
decodeQuality(quality, type = c("Illumina", "Sanger", "Solexa"))
encodeQuality(quality, type = c("Illumina", "Sanger", "Solexa"))
```

#### Arguments

- **quality**: For `decodeQuality` a character string representing the read qualities for a single sequence read. For `encodeQuality` a numeric vector of error probabilities.
- **type**: Type of encoding to use.

#### Details

See `extractQuality` for a description of the currently supported encodings.

#### Value

Either a numeric vector of error probabilities or a character string of encoded read quality scores. Each entry in the vector corresponds to one character in the input.

#### Author(s)

Peter Humburg

#### See Also

- `extractQuality`

#### Examples

```r
## decodeQuality and encodeQuality are the inverse operations
## of each other as one might expect
quality <- "IIIIIIIIIIIIIIIGII95II61II-II0"
errorProb <- decodeQuality(quality, type="Sanger")
qualitySanger <- encodeQuality(errorProb, type="Sanger")
all.equal(quality, qualitySanger)

## They can also be used to convert between encodings
qualityIllumina <- encodeQuality(errorProb, type="Illumina")
```
### defaultControl

**Default parameters for simChIP**

#### Description

Produces a list of parameters for each of the functions used to carry out the various stages of the simulation.

#### Usage

```r
defaultControl(features = list(), bindDensity = list(), readDensity = list(), readNames = list(), readSequence = list())
```

#### Arguments

- **features**: Parameters for feature generation.
- **bindDensity**: Parameters for the conversion of feature sequence into binding site densities.
- **readDensity**: Parameters for the conversion of binding site densities into read densities. Always provides parameters
  - `fragment` Default: `fragDens`
  - `meanLength` Default: 160
- **readNames**: Parameters for the generation of read names.
- **readSequence**: Parameters for the conversion of read positions into read sequences. Always provides parameters
  - `qualityFun` `readQualitySample`
  - `errorFun` `readError`
  - `readLen` 36

#### Details

Any parameters passed as part of list to one of the arguments of `defaultControl` will be passed on to the corresponding function in `simChIP`. The build-in defaults can be overwritten by providing a list entry with the same name.

#### Value

List of parameters for use as the `control` argument to `simChIP`.

#### Author(s)

Peter Humburg

#### See Also

`defaultFunctions`, `simChIP`

#### Examples

```r
defaultControl()
defaultControl(features=list(maxTail=0), readSequence=list(readLen=50))
```
defaultErrorProb  

*Replacement probabilities for sequencing errors*

**Description**

For each nucleotide this function provides probabilities indicating how likely it is to be replaced by any of the other nucleotides should a sequencing error occur.

**Usage**

```r
defaultErrorProb()
```

**Details**

The probabilities used here are the ones determined by Dohm et al. for *Beta vulgaris*. They should be more appropriate than a uniform distribution but the use of species specific rates is recommended where available.

**Value**

A list of four vectors with replacement probabilities for each nucleotide.

**Author(s)**

Peter Humburg

**References**


**Examples**

```r
defaultErrorProb()
```

---

**Default functions for simChIP**

**Description**

Provides default functions to carry out the different stages of the ChIP-seq simulation.

**Usage**

```r
defaultFunctions()
```
Value
A list with components

features  placeFeatures
bindDensity  feat2dens
readDensity  bindDens2readDens
sampleReads  sampleReads
readSequence  writeReads
readNames  simpleNames

Author(s)
Peter Humburg

See Also
simChIP

Examples

defaultFunctions()

---

Description
Functions to generate defaults for makeFeatures.

Usage
defaultGenerator()
defaultTransition()
defaultInit(prob=c(0.2, 0.05, 0, 0.25, 0.5), states=c("ReversePhasedFeature", "StableFeature", "PhasedFeature", "NFRFeature", "FuzzyFeature"))
defaultLastFeat(isEnd = c(FALSE, rep(TRUE, 4)), states = c("ReversePhasedFeature", "StableFeature", "PhasedFeature", "NFRFeature", "FuzzyFeature"))

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>prob</td>
<td>Numeric vector giving the initial state distribution. This will be normalised if the probabilities do not add up to 1.</td>
</tr>
<tr>
<td>isEnd</td>
<td>Logical vector indicating which states, i.e. features, are allowed to be last in the sequence.</td>
</tr>
<tr>
<td>states</td>
<td>Character vector of state names.</td>
</tr>
</tbody>
</table>
extractQuality

Details

These functions generate data structures that can be passed as arguments to `makeFeatures`. Using this set of functions will create a nucleosome positioning simulation. Some of the defaults can be modified by passing different values to `defaultInit` and `defaultLastFeat`.

Value

Return values are suitable as arguments `generator`, `transition`, `init` and `lastFeat` of `makeFeatures`. See the documentation of `makeFeatures` for more detail.

Author(s)

Peter Humburg

Examples

```r
set.seed(1)
## generate defaults
generator <- defaultGenerator()
transition <- defaultTransition()
lastFeat <- defaultLastFeat()

## change the initial state distribution such that it
## always starts with a fuzzy feature
init <- defaultInit(c(0, 0, 0, 0, 1))

## now generate some features for a stretch of 1 million base pairs
features <- makeFeatures(generator=generator, transition=transition,
lastFeat=lastFeat, init=init, length=1e6)
```

extractQuality

Obtain read qualities from a Fastq file or ShortReadQ object

Description

Converts the read qualities encoded in fastq formatted files into error probabilities.

Usage

```r
extractQuality(reads, minLength = 25, dir,
type = c("Illumina", "Sanger", "Solexa"))
```

Arguments

- `reads` Either the name of a fastq file or a `ShortReadQ` object (see Details).
- `minLength` Minimum read length required.
- `dir` Directory of fastq file.
- `type` Character string indicating the format the qualities are encoded in (see Details).
Details

If `reads` and `dir` are character strings it is assumed that `dir/reads` is the name of a fastq file. Otherwise `reads` should be a `ShortReadQ` object in which case `dir` is ignored.

Currently three different encodings of read qualities are supported. The encoding has to be selected via the `type` argument. The supported formats are

**Illumina** The format currently used by Illumina (version 1.3). This is a phred score between 0 and 40 encoded as ASCII characters 64 to 104. [default]

**Sanger** The Sanger format uses a phred quality score between 0 and 93 encoded as ASCII characters 33 to 126.

**Solexa** The old Solexa format previously used by Solexa/Illumina uses a quality score between -5 and 40 encoded as ASCII characters 59 to 104.

Value

A list with a vector of error probabilities for each read in `reads` that is at least `minLength` nucleotides long.

Author(s)

Peter Humburg

See Also

`decodeQuality`, `readQualitySample`

Examples

```r
## Not run:
## load reads from a fastq file with Sanger encoding
qualities <- extractQuality("test.fastq", dir=".", type="Sanger")

## extract error probabilities for first 25bp of each read
qualities25 <- sapply(qualities, "[", 1:25)

## plot average quality for each position
plot(rowMeans(qualities25), type='b', xlab="Read position", ylab="Error probability")
## End(Not run)
```

---

**feat2dens**

*Convert a list of features into a feature density*

**Description**

Given a list of features (as produced by `makeFeatures`) computes the feature density for each and combines them into a chromosome wide density.

**Usage**

`feat2dens(features, length, featureBgr = TRUE, ...)`
**Arguments**

- **features**
  A list of features.

- **length**
  Total length of feature density vector (i.e. chromosome length). If this is missing
  the length is inferred from the feature parameters.

- **featureBgr**
  Logical indicating whether feature specific background should be added to the
  density. If this is **TRUE** the resulting density for each feature is a mixture of the
  feature density and a fuzzy, i.e. uniform, feature density. The weights of the
  components are determined by the feature weight.

... Further arguments to `featureDensity`.

**Value**

A vector with the feature density for each position along the chromosome.

**Author(s)**

Peter Humburg

**See Also**

The majority of the work is done by calls to `featureDensity` and `joinRegion`.

**Examples**

```r
set.seed(1)
## generate a (relatively short) sequence of nucleosome features
features <- placeFeatures(start=200, length=1e5)
## calculate density
featureDens <- feat2dens(features, length=1e5)
```

**Description**

This set of functions is used to generate the density of individual features of different types. `featureDensity` is an S3 generic, functions may be defined for different feature classes.

**Usage**

```r
featureDensity(x, ...)
## S3 method for class 'StableFeature'
featureDensity(x, stable=stableDens, background=FALSE, ...)
## S3 method for class 'StablePhasedFeature'
featureDensity(x, stable=stableDens, dist=distDens, background=FALSE, ...)
## S3 method for class 'ReversePhasedFeature'
featureDensity(x, stable=stableDens, dist=distDens, background=FALSE, ...)
## S3 method for class 'NFRFeature'
featureDensity(x, background=FALSE, ...)
## S3 method for class 'FuzzyFeature'
featureDensity(x, ...)
```
Arguments

- **x**: The feature for which the density should be computed.
- **stable**: Function that should be used to compute the density of a stable feature.
- **dist**: Function that should be used to compute the distribution of distances between adjacent features.
- **background**: Logical indicating whether uniform background should be added to the feature.
- **...**: Arguments to future functions.

Details

These functions are used internally by `feat2dens`. There should be no need to call them directly but it is important to supply suitable `featureDensity` functions for new feature types.

Value

A two column matrix. The first column contains the density, the second the weight at each position.

Author(s)

Peter Humbug

See Also

`feat2dens`, `makeFeatures`

Examples

```r
## Create a single StableFeature
feature <- stableFeature(start = 200, weight = 0.8, shift = 10, stability = 1, ratio = 1)

## Convert the feature into a density (without background)
featDens <- featureDensity(feature)

## If we want featureDensity to automatically add uniform background
## we have to ensure that the feature has a 'meanDist' parameter
## (this is usually added by 'reconcileFeatures').
feature$meanDist <- 200
featDens2 <- featureDensity(feature, background = TRUE)
```

Description

These functions are used to generate the parameters for different nucleosome positioning related features.
Usage

stableFeature(start, minDist = 175, weight = seq(0.1, 1, by = 0.1),
shift = c(0, 5, 10), ratio = seq(0, 4, by = 0.25),
stability = seq(0.1, 5, by = 0.1), weightProb, shiftProb,
ratioProb, stabilityProb, ...)
phasedFeature(minDist = 175, length = seq(1000, 10000, by = minDist),
meanDist = minDist:300, lengthProb, meanDistProb, start, ...)
fuzzyFeature(start, length = seq(1000, 10000, by = 1000),
meanDist = 175:400, lengthProb, meanDistProb, ...)
nfrFeature(start, length = seq(50, 500, by = 10),
weight = seq(0.1, 1, by = 0.1), lengthProb, weightProb, ...)

Arguments

- **start**: Start location of feature on chromosome.
- **minDist**: Minimum distance between nucleosomes.
- **length**: A numeric vector giving possible values for the length of the feature.
- **meanDist**: A numeric vector giving possible values for the mean distance between nucleosomes.
- **weight**: A numeric vector giving possible values for the weight of the feature.
- **shift**: A numeric vector giving possible values for the distance between favoured positions of stable nucleosomes.
- **ratio**: A numeric vector giving possible values for the ratio between probabilities for alternative and central position of stable nucleosomes.
- **stability**: A numeric vector giving possible values for the stability of stable nucleosomes.
- **lengthProb**: Length distribution of feature. This corresponds to the state duration distribution of the underlying generating model.
- **meanDistProb**: Distribution of mean distances between nucleosomes.
- **weightProb**: Distribution of feature weights.
- **shiftProb**: Distribution of distances between favoured positions of stable nucleosome.
- **ratioProb**: Ratio distribution.
- **stabilityProb**: Stability distribution.
- **...**: Further arguments (currently ignored).

Value

A list of parameters for the corresponding feature type. These parameters are later used to compute nucleosome densities.

Author(s)

Peter Humburg

See Also

- `simChIP`

Examples

```R
feature <- stableFeature(200)
```
makeFeatures

Generating a list of genomic features

Description

This function generates a list of genomic features for a single chromosome based on a Markov model.

Usage

makeFeatures(generator = defaultGenerator(),
transition = defaultTransition(), init = defaultInit(),
start = 1000, length, control = list(),
globals = list(minDist = 175), lastFeat = defaultLastFeat(),
experimentType = "NucleosomePosition",
truncate = FALSE, maxTries = 10, force=FALSE)

Arguments

generator A named list providing functions to generate the parameters associated with each type of feature. The name of each element in the list is the name of the state the function should be associated with.

transition Named list of transition probabilities. Each element is a (named) numeric vector giving the transition probabilities for the state indicated by the element’s name, i.e., each element of the list is a row of the transition probability matrix but zero probabilities can be omitted.

init Named numeric vector of initial state probabilities. The names have to correspond to state names of the model. Zero probabilities may be omitted.

start Numeric value indicating the position at which the first feature should be placed.

length Maximum length of DNA that should be covered with features.

control Named list with additional arguments to generator functions (one list per generator). Again the names should be the same as the state names.

globals List of global arguments to be passed to all generator functions.

lastFeat Named logical vector indicating for each feature type whether it can be the last feature.

experimentType Type of experiment the simulated features belong to. This is used as the class of the return value.

truncate Logical value indicating whether the final feature should be truncated to ensure that total length does not exceed length (if FALSE, a feature that would be truncated is removed instead).

maxTries Maximum number of attempts made to generate a valid sequence of features. If no valid sequence is generated during the first maxTries attempts the function will fail either silently (returning an empty sequence) or raise an error, depending on the value of force.

force Logical indicating whether the function should be forced to return a feature sequence, even if no valid sequence was found. If this is TRUE an empty sequence will be returned in that case otherwise an error is raised.
makeFeatures

Details

This function will generate features from any first order Markov model specified by init, transition and generator. If force is FALSE the returned feature sequence is guaranteed to contain at least one feature and end in a state that is indicated as possible end state in lastFeat. Note that the states for which lastFeat is TRUE are not end states in the sense that the chain is terminated once the state is entered or that the chain remains in the state once it is first entered. Instead this is a mechanism to ensure that some states cannot be the last in the sequence.

Due to the constrains on the total length of DNA covered by features as well as the possible constraint on the final feature of the sequence it is possible to specify models that cannot produce a legal sequence. In other cases it may be possible but unlikely to produce a feature sequence that satisfies both constraints. A serious attempt is made to satisfy both requirement, generating a new feature sequence or truncating an existing one if necessary. To ensure that this terminates eventually the number of attempts to generate a valid sequence are limited to maxTries.

In some cases it may be desirable to carry out some post-processing of the feature sequence to ensure that parameters of neighbouring features are compatible in some sense. For the default nucleosome positioning simulation the function reconcileFeatures provides this functionality and placeFeatures is an interface to makeFeatures that utilises reconcileFeatures.

Value

A list of features (with class determined by experimentType). Each feature is represented by a list of parameters and has a class with the same name as the state that generated the feature. In addition all features are of class SimulatedFeature.

Author(s)

Peter Humburg

See Also

Functions to generate default values for some of the arguments: defaultGenerator, defaultInit, defaultTransition, defaultLastFeat.

Use feat2dens to convert a feature sequence into feature densities. placeFeatures is an interface to makeFeature for nucleosome positioning.

Examples

```r
set.seed(1)
## generate a (relatively short) sequence of nucleosome features
features <- makeFeatures(length=1e6)

## check the total length of the features
sum(sapply(features, "[[", "length")) # 995020
```
placeFeatures

Generating and reconciling a feature sequence

Description

This function provides an interface to makeFeatures and reconcileFeatures that combines both steps of the feature generation process.

Usage

placeFeatures(..., maxTail = 0.01, compoundFeatures=list("StablePhasedFeature"))

Arguments

... Arguments to makeFeatures.
maxTail Maximum portion of total length of chromosome that may be left free of features (see Details).
compoundFeatures List of feature classes that are produced by combining two features. This may happen during the call to reconcileFeatures and requires special handling when extending the feature list.

Details

This function (as well as makeFeatures which it calls) tries to fill as much of the genomic region with features as possible, i.e. an attempt is made to produce a feature sequence that covers length base pairs. In most cases the sequence will be slightly shorter. The maxTail argument determines how long a region without any features at the end of the genomic region is acceptable (as fraction of the total length). Note however that even maxTail = 0 does not guarantee a feature sequence of exactly the requested length.

Value

A list of simulated features. The class of the return value as well as the features generated depend on the arguments passed to makeFeatures.

Note

Using the reconcileFeatures mechanism it is possible to introduce dependence between neighbouring features that is not easily expressed in terms of a simple Markov model. In some cases the same effect can be achieved by introducing additional states into the model but it may be more convenient to simply post-process the feature sequence.

Author(s)

Peter Humburg

See Also

makeFeatures, reconcileFeatures
Examples

```r
set.seed(1)
## generate a (relatively short) sequence of nucleosome features
features <- placeFeatures(length=1e6, maxTail = 0)

## check the total length of the features
sum(sapply(features, "[[", "length")) ## 990509
```

pos2fastq

Convert read positions to fastq records

Description

Convert read positions for a single chromosome (both strands) into read sequences + qualities and write them to file

Usage

```r
pos2fastq(readPos, names, quality, sequence, qualityFun, errorFun, 
readLen = 36, file, 
qualityType = c("Illumina", "Sanger", "Solexa"), ...)
```

Arguments

- **readPos**: A list of two numeric vectors (one per strand)
- **names**: List of names to use for reads in fastq file. Has to be of same shape as name.
- **quality**: Passed on as argument to qualityFun.
- **sequence**: Reference sequence (a DNAString object).
- **qualityFun**: Function to generate quality scores.
- **errorFun**: Function to introduce sequencing errors.
- **readLen**: Read length to generate.
- **file**: Output file (either file name or connection).
- **qualityType**: Encoding to use for read quality scores.
- **...**: Further arguments (see Details).

Details

Arguments passed as part of ... will be passed on to qualityFun, except an argument called prob which is passed on to errorFun instead if present.

Value

Invisibly returns the number of records that were written.

Author(s)

Peter Humburg
See Also

See `readError` for a possible choice of `errorFun` and `readQualitySample` for a simple `qualityFun`.

Examples

```r
set.seed(1)

## a function to generate random read qualities (in Sanger format)
randomQuality <- function(read, ...){
paste(sample(unlist(strsplit(rawToChar(as.raw(33:126)),"")),
length(read), replace = TRUE), collapse="")
}

## generate a reference sequence
chromosome <- DNAString(paste(sample(c("A", "C", "G", "T"),
1e5, replace = TRUE), collapse = ""))

## and a few read positions
reads <- list(sample(100:9900, 5), sample(100:9900, 5))
names <- list(paste("read", 1:5, sep="_"), paste("read", 6:10, sep="_"))

## convert to fastq format
pos2fastq(reads, names, sequence=chromosome, qualityFun=randomQuality,
errorFun=readError, file="")
```

---

### readError

**Introduce errors into read sequence based on quality scores**

**Description**

Given a read sequence and quality this function introduces errors by first choosing positions that should be modified based on the quality score and then exchanges nucleotides based on the probabilities given in `prob`.

**Usage**

```r
readError(read, qual, alphabet = c("A", "C", "G", "T"),
prob = defaultErrorProb(), ...)
```

**Arguments**

- `read`: A character string representing a read sequence.
- `qual`: Numeric vector of read error probabilities.
- `alphabet`: Alphabet used for read sequence.
- `prob`: Nucleotide exchange probabilities.
- `...`: Further arguments (currently ignored).

**Details**

If the read sequence contains letters that are not part of `alphabet` they are replaced by the first entry of `alphabet` before positions of sequencing errors are determined. The alphabet used has to match the names used in `prob`.
Value

The modified read sequence.

Author(s)

Peter Humburg

See Also

defaultErrorProb, readSequence

Examples

```r
set.seed(42)

## generate sequence read and quality
quality <- paste(sample(unlist(strsplit(rawToChar(as.raw(33:126)),"")),
36, replace = TRUE), collapse="")
errorProb <- decodeQuality(quality, type = "Sanger")
read <- paste(sample(c("A", "C", "G", "T"), 36, replace = TRUE),
collapse = "")

## use readError to introduce sequencing errors
read2 <- readError(read, errorProb)

all.equal(read, read2) ## "1 string mismatch"
```

Description

Given a read sequence and a list of read quality scores this function returns a (possibly truncated) quality score of the same length as the read.

Usage

```r
readQualitySample(read, qualities, checkLength = TRUE, ...)
```

Arguments

- `read`: A sequence read.
- `qualities`: List of sequence read quality scores.
- `checkLength`: Flag indicating whether the length of quality scores should be checked to ensure that they are at least as long as the read. If `qualities` contains entries shorter than `read` this has to be `TRUE`, but see below.
- `...`: Further arguments, currently not used.

Details

Using `checkLength = TRUE` leads to a substantial decrease in performance and is impractical for a large simulation. To avoid this slow down it is recommended to remove short sequences from `qualities` beforehand so that `checkLength = FALSE` can be used.
readSequence

Value

An read quality score string of the same length as read.

Author(s)

Peter Humburg

---

**readSequence**  
*Convert read position into read sequence*

---

**Description**

Given a read position, a reference sequence, a strand and a read length this function returns the read sequence.

**Usage**

```
readSequence(readPos, sequence, strand, readLen = 36)
```

**Arguments**

- `readPos` Numeric value indicating the start position on the chromosome.
- `sequence` Chromosome sequence (a DNAString)
- `strand` Strand indicator (+1 /-1)
- `readLen` Length of read.

**Value**

Read sequence.

**Author(s)**

Peter Humburg

**See Also**

`readError`, `writeReads`
**reconcileFeatures**  
*Post-processing of simulated features*

**Description**  
The `reconcileFeatures` functions provide a facility to post-process a list of features representing a simulated experiment. `reconcileFeatures` is an S3 generic, new functions can be added for additional types of experiment. The current default is to call `reconcileFeatures.SimulatedExperiment` which, if called without further arguments, will simply return the feature list unchanged.

**Usage**  
```r  
reconcileFeatures(features, ...)  
## Default S3 method:  
reconcileFeatures(features, ...)  
## S3 method for class 'SimulatedExperiment'  
reconcileFeatures(features, defaultValues=list(), ...)  
## S3 method for class 'NucleosomePosition'  
reconcileFeatures(features, defaultMeanDist = 200, ...)  
```

**Arguments**  
- `features`  
  List of simulated features.
- `defaultValues`  
  Named list of default parameter values. The method for class `SimulatedExperiment` ensures that all features have at least the parameters listed in `defaultValues`, adding them where necessary.
- `defaultMeanDist`  
  Default value for the average distance between nucleosomes for nucleosome positioning experiments.
- `...`  
  Further arguments to future functions.

**Value**  
A list of features of the same class as `features`.

**Author(s)**  
Peter Humburg

**See Also**  
`makeFeatures`, `placeFeatures`

**Examples**  
```r  
set.seed(1)  
## generate a (relatively short) sequence of nucleosome features  
features <- makeFeatures(length=1e6)  
## check the total length of the features  
sum(sapply(features, "[[", "length"))  
## 995020  
```
sampleReads

## reconcile features to ensure smooth transitions
## For experiments of class NucleosomePosition this
## also combines some features and introduces
## some overlap between them.
features <- reconcileFeatures(features)

## check the total length of the features again
sum(sapply(features, function(x) length(x))) # 984170

---

### sampleReads

*Sampling sequence read positions from a read density.*

**Description**

Given a read density this function returns the starting positions of sequence reads.

**Usage**

```r
sampleReads(readDens, nreads = 6e+06, strandProb = c(0.5, 0.5))
```

**Arguments**

- `readDens`: A two column matrix of read densities (as produced by `bindDens2readDens`).
- `nreads`: Number of read positions to generate.
- `strandProb`: A numeric vector with two elements giving weights for forward and reverse strand.

**Details**

The expected number of reads for each strand is `strandProb * nreads`.

**Value**

A list with components `fwd` and `rev` giving the read positions on the forward and reverse strand respectively.

**Author(s)**

Peter Humburg

**See Also**

`bindDens2readDens`
Examples

```r
caseRseed(1)
## generate a (relatively short) sequence of nucleosome features
features <- placeFeatures(start=200, length=1e5)

## calculate feature density
featureDens <- feat2dens(features, length=1e5)

## convert to read density
readDens <- bindDens2readDens(featureDens, fragDens, meanLength=160)

## sample reads
## of course you would usually want a much larger number
readPos <- sampleReads(readDens, nreads=1000)
```

---

**simChIP**

*Simulate ChIP-seq experiments*

**Description**

This function acts as driver for the simulation. It takes all required arguments and passes them on to the functions for the different stages of the simulation. The current defaults will simulate a nucleosome positioning experiment.

**Usage**

```r
simChIP(nreads, genome, file, functions = defaultFunctions(),
control = defaultControl(), verbose = TRUE, load = FALSE)
```

**Arguments**

- `nreads`: Number of reads to generate.
- `genome`: An object of class ‘DNAStringSet’ or the name of a fasta file containing the genome sequence.
- `file`: Base of output file names (see Details).
- `control`: Named list of arguments to be passed to simulation functions (one list per function).
- `verbose`: Logical indicating whether progress messages should be printed.
- `load`: Logical indicating whether an attempt should be made to load intermediate results from a previous run.
Details

The simulation consists of six stages:

1. generate feature sequence (for each chromosome): chromosome length -> feature sequence (list)
2. compute binding site density: feature sequence -> binding site density (vector)
3. compute read density: binding site density -> read density (two column matrix, one column for each strand)
4. sample read start sites: read density -> read positions (list)
5. create read names: number of reads -> unique names
6. obtain read sequence and quality: read positions, genome sequence, [qualities] -> output file

After each of the first three stages the results of the stage are written to a file and can be reused later. File names are created by appending ‘_features.rdata’, ‘_bindDensity.rdata’ and ‘_readDensity.rdata’ to file respectively. Previous results will be loaded for reuse if load is TRUE and files with matching names are found. This is useful to sample repeatedly from the same read density or to recover partial results from an interrupted run.

The creation of files can be prevented by setting file = “”. In this case all results will be returned in a list at the end. Note that this will require more memory since all intermediate results have to be held until the end.

The behaviour of the simulation is mainly controlled through the functions and control arguments. They are expected to be lists of the same length with matching names. The names indicate the stage of the simulation for which the function should be used; elements of control will be used as arguments for the corresponding functions.

Value

A list. The components are typically either lists (with one component per chromosome) or file names but note that this may depend on the return value of functions listed in functions. The components of the returned list are:

- `features` Either a list of generated features or the name of a file containing that list;
- `bindDensity` Either a list with binding site densities or the name of a file containing that list;
- `readDensity` Either a list of read densities or the name of a file containing that list;
- `readPosition` Either a list of read start sites or the name of a file containing that list;
- `readSequence` The return value of the function listed as ‘readSequence’. The default for this the name of the fastq file containing the read sequences;
- `readNames` Either a list of read names or the name of a file containing that list.

Author(s)

Peter Humburg

See Also

defaultFunctions, defaultControl
Examples

## Not run:
## To run the default nucleosome positioning simulation
## we can simply run something like the line below.
## This will result in 10 million reads sampled from the genome.
## Of course the file names have to be changed as appropriate.
simChIP(1e7, genome = "reference.fasta", file = "output/sim_10M")

## End(Not run)

---

simpleNames  Generate unique read names

Description

Generates a set of unique (and very simple) read names.

Usage

```r
simpleNames(n, nameBase = "read")
```

Arguments

- `n`: Number of names to generate.
- `nameBase`: Base name to use.

Value

A character vector with entries of the form ‘nameBase_i’ where i runs from 1 to n.

Author(s)

Peter Humburg

Examples

```r
simpleNames(5)
```

---

writeFASTQ  Write read sequences and qualities to a FASTQ formatted file

Description

This is intended to produce the final output of the simulation by providing a fastq file that may then be used in an analysis pipeline.

Usage

```r
writeFASTQ(read, quality, name, file, append = FALSE)
```
writeReads

Arguments

- **read**: List of read sequences.
- **quality**: List of read quality scores.
- **name**: Read names.
- **file**: File name. If this is "" results will be printed to the standard output connection.
- **append**: Logical indicating the reads should be appended to an existing file.

Details

The first three arguments should have the same length but will be recycled if necessary.

Value

Called for its side effect.

Author(s)

Peter Humburg

See Also

readSequence, readQualitySample, writeReads

Examples

```r
set.seed(1)

## generate sequence read and quality
quality <- paste(sample(unlist(strsplit(rawToChar(as.raw(33:126)),"")), 36, replace = TRUE), collapse="")
read <- paste(sample(c("A", "C", "G", "T"), 36, replace = TRUE), collapse = "")

## write a fastq record
writeFASTQ(read, quality, "read_1", file="")
```

writeReads

Create fastq file from read positions

Description

This is an interface to pos2fastq that writes all reads for a given genome to a single file.

Usage

writeReads(readPos, readNames, sequence, quality, file, ...)

Arguments

- **readPos**
  List of read positions with each component holding the read positions for one chromosome, which are themselves two component lists that provide forward and reverse strand positions.

- **readNames**
  List of the same shape as `readPos` providing read names.

- **sequence**
  Genome reference sequence (a `DNAStringSet`).

- **quality**
  Read quality scores (see Details).

- **file**
  Output file.

- **...**
  Further arguments to `pos2fastq`.

Details

If `quality` looks like it might refer to a fastq file an attempt is made to create a `ShortReadQ` object. The read qualities of any `ShortReadQ` object passed as `quality` (directly or indirectly as file name) are extracted and passed on to `pos2fastq` as `quality` argument. Otherwise it is passed on unchanged.

Value

The name of the output file.

Author(s)

Peter Humburg

See Also

- `pos2fastq`
Index

*Topic datagen
bindDens2readDens, 3
defaultGenerator, 7
feat2dens, 9
featureDensity, 10
FeatureGenerators, 11
makeFeatures, 13
placeFeatures, 15
readError, 17
readQualitySample, 18
reconcileFeatures, 20
sampleReads, 21
simChIP, 22
*Topic package
ChIPsim-package, 2
*Topic utilities
decodeQuality, 4
defaultControl, 5
defaultErrorProb, 6
defaultFunctions, 6
defaultGenerator, 7
extractQuality, 8
pos2fastq, 16
readQualitySample, 18
readSequence, 19
simpleNames, 24
writeFASTQ, 24
writeReads, 25
bindDens2readDens, 3, 7, 21
ChIPsim (ChIPsim-package), 2
ChIPsim-package, 2
defaultTransition (defaultGenerator), 7
DNAString, 16, 19
DNAStringSet, 26
encodeQuality (decodeQuality), 4
extractQuality, 4, 8
feat2dens, 3, 7, 9, 11, 14
featureDensity, 10, 10
FeatureGenerators, 11
fragDens, 5
fuzzyFeature (FeatureGenerators), 11
joinRegion, 10
makeFeatures, 7–9, 11, 13, 15, 20
nfrFeature (FeatureGenerators), 11
phasedFeature (FeatureGenerators), 11
placeFeatures, 7, 14, 15, 20
pos2fastq, 16, 25, 26
readError, 5, 17, 17, 19
readQualitySample, 5, 9, 17, 18, 25
readSequence, 18, 19, 25
reconcileFeatures, 14, 15, 20
sampleReads, 3, 7, 21
ShortRead, 2
ShortReadQ, 8, 9, 26
simChIP, 2, 5, 7, 12, 22
simpleNames, 7, 24
stableFeature (FeatureGenerators), 11
writeFASTQ, 24
writeReads, 7, 19, 25, 25