# Package 'cleanUpdTSeq'

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Type Package

**Title** cleanUpdTSeq cleans up artifacts from polyadenylation sites from oligo(dT)-mediated 3' end RNA sequending data

Description This package implements a Naive Bayes classifier for accurately differentiating true polyadenylation sites (pA sites) from oligo(dT)-mediated 3' end sequencing such as PAS-Seq, PolyA-Seq and RNA-Seq by filtering out false polyadenylation sites, mainly due to oligo(dT)-mediated internal priming during reverse transcription. The classifer is highly accurate and outperforms other heuristic methods.

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**Depends** R (>= 3.5.0), BSgenome.Drerio.UCSC.danRer7, methods

**Imports** BSgenome, GenomicRanges, seqinr, e1071, Biostrings, GenomeInfoDb, IRanges, utils, stringr, stats, S4Vectors

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BED6WithSeq2GRangesSeq

Covert (extended) BED6 file to a GRanges object

# **Description**

Convert to a GRanges object from a (extended) BED6 file with at least six columns: chrom, chrom-Start, strEnd, name, score and strand, and optional upstream sequences (including pA sites) and downstream sequences of pA sites

# Usage

```
BED6WithSeq2GRangesSeq(
   file,
   skip = 1L,
   withSeq = TRUE,
   upstream.seq.ind = 7L,
   downstream.seq.ind = 8L
)
```

# **Arguments**

A character(1) vector, representing a path to a extended BED file containing at least six columns in the order of chrom, chromStart, strEnd, name, score and strand. The strand information must be designated as "+", or "-". Optional fields—upstream sequences (including pA sites) and downstream sequences of pA sites—are allowed. For more details about the BED format, see https://genome.ucsc.edu/FAQ/FAQ/Skip A integer(1) vector, indicating how many rows (header lines) to skip when the BED file is read into R.

withSeq A logical(1) vector, indicating that upstream and downstream sequences flank-

upstream.seq.ind

An integer(1),vector delineating the column location of upstream sequences of the putative pA site

ing pA sites are included in the file

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```
downstream.seq.ind
```

An integer(1),vector delineating the column location of downstream sequences of the putative pA site

### Value

An object of GRanges

### Author(s)

Haibo Liu, Lihua J. Zhu

### **Examples**

buildClassifier

Build a Naive Bayes Classifier

# Description

Computes the conditional a-posterior probabilities of a categorical class variable given independent predictor variables using the Bayes rule.

## Usage

```
buildClassifier(
  Ndata.NaiveBayes,
  Pdata.NaiveBayes,
  upstream = 40L,
  downstream = 30L,
  wordSize = 6L,
  alphabet = c("ACGT")
)
```

# Arguments

Ndata.NaiveBayes

A data.frame, containing features for the negative training data, described further in data.NaiveBayes.

Pdata.NaiveBayes

A data.frame, containing features for the positive training data, described further in data.NaiveBayes.

upstream An integer(1) vector, length of upstream sequence to retrieve.

An integer(1) vector, length of downstream sequence to retrieve.

wordSize An integer(1) vector, size of the kmer feature for the upstream sequence. word-

Size = 6 should always be used.

alphabet A character(1) vector, a string containing DNA bases. By default, "ACTG".

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#### Value

An object of class "naiveBayes".

### Author(s)

Jianhong Ou

### See Also

naiveBayes

# **Examples**

buildFeatureVector

build Feature Vector\_2

# **Description**

This function creates a data frame. Fields include peak name, upstream sequence, downstream sequence, and features to be used in classifying the putative polyadenylation site.

# Usage

```
buildFeatureVector(
  peaks,
  genome = Drerio,
  upstream = 40L,
  downstream = 30L,
  wordSize = 6L,
  alphabet = "ACGT",
  sampleType = c("TP", "TN", "unknown"),
  replaceNAdistance = 30L,
  method = c("NaiveBayes", "SVM"),
  fetchSeq = FALSE,
  return_sequences = FALSE
)
```

# **Arguments**

peaks An object of GRanges that may contain the upstream and downstream sequence

information. This item is created by the function BED6WithSeq2GRangesSeq.

genome Name of the genome to get sequences from. To find out a list of available

genomes, please type BSgenome::available.genomes() in R.

upstream An integer(1) vector, length of upstream sequence to retrieve.

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downstream An integer(1) vector, length of downstream sequence to retrieve.

wordSize An integer(1) vector, size of the kmer feature for the upstream sequence. word-

Size = 6 should always be used.

alphabet A character(1) vector, a string containing DNA bases. By default, "ACTG".

sampleType A character(1) vector, indicating type of sequences for building feature vectors.

Options are TP (true positive) and TN (true negative) for training data, or un-

known for test data.

replaceNAdistance

An integer(1) vector, specifying an number for avg.distanceA2PeakEnd, the average distance of As to the putative pA site, when there is no A in the down-

stream sequence.

method A character(1) vector, specifying a machine learning method to use. Cur-

rently, only "NaiveBayes" is implemented.

fetchSeq A logical (1), indicating whether upstream and downstream sequences should

be retrieved from the BSgenome object at this step or not.

return\_sequences

A logical(1) vector, indicating whether upstream and downstream sequences

should be included in the output

### Value

An object of "featureVector"

#### Author(s)

Sarah Sheppard, Haibo Liu, Jianhong Ou, Nathan Lawson, Lihua J. Zhu

```
library(BSgenome.Drerio.UCSC.danRer7)
testFile <- system.file("extdata", "test.bed",</pre>
                        package = "cleanUpdTSeq")
peaks <- BED6WithSeq2GRangesSeq(file = testFile,</pre>
                                skip = 1L, withSeq = TRUE)
## build the feature vector for the test set with sequence information
testSet.NaiveBayes = buildFeatureVector(peaks,
                                         genome = Drerio,
                                         upstream = 40L,
                                         downstream = 30L,
                                         wordSize = 6L,
                                         alphabet = "ACGT",
                                         sampleType = "unknown";
                                         replaceNAdistance = 30,
                                         method = "NaiveBayes",
                                         fetchSeq = FALSE,
                                         return_sequences = TRUE)
## convert the test set to GRanges without upstream and downstream
## sequence information
peaks <- BED6WithSeq2GRangesSeq(file = testFile,</pre>
                                skip = 1L, withSeq = FALSE)
#build the feature vector for the test set without sequence information
testSet.NaiveBayes = buildFeatureVector(peaks,
                                         genome = Drerio,
```

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```
upstream = 40L,
downstream = 30L,
wordSize = 6L,
alphabet = "ACGT",
sampleType = "unknown",
replaceNAdistance = 30,
method = "NaiveBayes",
fetchSeq = TRUE,
return_sequences = TRUE)
```

classifier

NaiveBayes classifier

# Description

An object of class "naiveBayes" generated from data. NaiveBayes

# Usage

classifier

#### **Format**

An object of class "PASclassifier" including components:

# **Examples**

```
data(classifier)
names(classifier)
```

cleanUpdTSeq

This package classifies putative polyadenylation sites.

# **Description**

3'ends of transcripts have generally been poorly annotated. With the advent of deep sequencing, many methods have been developed to identify 3'ends. The majority of these methods use an oligodT primer which can bind to internal adenine-rich sequences, and lead to artifactual identification of polyadenylation sites. Heuristic filtering methods rely on a certain number of As downstream of a putative polyadenylation site to classify the site as true or oligodT primed. This package provides a robust method to classify putative polyadenylation sites using a Naive Bayes classifier.

# Author(s)

Sarah Sheppard, Haibo Liu, Jianhong Ou, Nathan Lawson, Lihua Julie Zhu

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data.NaiveBayes

Training Data

# Description

A RData containing negative and positive training data

# Usage

data.NaiveBayes

#### **Format**

A list with 2 data frame, "Negative" and "Positive". Negative has 9219 observations on the following 4120 variables. And Positive is a data frame with 22770 observations on the following 4120 variables. The format is:

```
list("Negative") 'data.frame': 9219 obs. of 4120 variables:
```

list("Positive") 'data.frame': 22770 obs. of 4120 variables:

Both of them have same structure.

list("y") a numeric vector

list("n.A.Downstream") a numeric vector

list("n.C.Downstream") a numeric vector

list("n.T.Downstream") a numeric vector

list("n.G.Downstream") a numeric vector

list("avg.distanceA2PeakEnd") a numeric vector

list("dimer") a numeric vector

: such as AA, AC, AG, AT, CA, ... etc. a numeric vector

list("heximer") a factor with levels 0 1

: such as AAAAAA, ACGTAC, ... etc. a factor with levels 0 1

list("upstream.seq") a vector of sequence string

list("downstream.seq") a vector of sequence string

```
library(BSgenome.Drerio.UCSC.danRer7)
data(data.NaiveBayes)
head(str(data.NaiveBayes$Negative))
head(str(data.NaiveBayes$Positive))
```

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```
featureVector-class Class "featureVector"
```

#### **Description**

An object of class "featureVector" represents the output of buildFeatureVector

# **Objects from the Class**

Objects can be created by calls of the form new("featureVector", data, info).

getContextSequences Retrieve upstream and downstream sequences

### **Description**

Retrieve upstream and downstream sequences of pA sites from a BSgenome object based on a GRanges object

## Usage

```
getContextSequences(peaks, upstream = 40L, downstream = 30L, genome)
```

### Arguments

peaks An object of GRanges representing pA sites

upstream An integer(1) vector, length of upstream sequence of pA sites, including pA site.

downstream An integer(1) vector, length of downstream sequences of pA sites

genome An object of BSgenome.

# Value

A data.frame containing sequences upstream and downstream pA sites:

```
upstream.seq sequence upstream pA site, including pA site downstream.seq sequence downstream pA site
```

### Author(s)

Haibo Liu

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modelInfo-class	Class "modelInfo"

### **Description**

An object of class "modelInfo" represents the information of sequence to use in the analysis

# **Objects from the Class**

Objects can be created by calls of the form new("modelInfo", upstream, downstream, wordSize, alphabet).

naiveBayes-class Class "naiveBayes"

# **Description**

An object of class "naiveBayes" represents the conditional a-posterior probabilities of a categorical class variable given independent predictor variables using the Bayes rule.

# Objects from the Class

Objects can be created by calls of the form new("naiveBayes", apriori, tables, levels, call).

PASclassifier-class Class "PASclassifier"

# **Description**

An object of class "PASclassifier" represents the output of buildClassifier

# **Objects from the Class**

Objects can be created by calls of the form new("PASclassifier", classifier, info).

### **Examples**

data(classifier)
classifier\$info\$upstream
classifier\$info\$wordSize
classifier\$info\$alphabet

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predictTestSet

predict authenticity of putative pA sites

### **Description**

classify putative pA sites into true and false bins.

#### Usage

```
predictTestSet(
  Ndata.NaiveBayes = NULL,
  Pdata.NaiveBayes = NULL,
  testSet.NaiveBayes,
  classifier = NULL,
  outputFile = "test-predNaiveBayes.tsv",
  assignmentCutoff = 0.5,
  return_sequences = FALSE
)
```

#### **Arguments**

Ndata.NaiveBayes

A data.frame, containing features for the negative training data, which is built using the function buildFeatureVector. It is described further indata.NaiveBayes.

Pdata.NaiveBayes

A data.frame, containing features for the positive training data, which is built using the function buildFeatureVector. It is described further indata.NaiveBayes.

testSet.NaiveBayes

An object of featureVector for test data built for Naive Bayes analysis using the function buildFeatureVector.

An object of class PASclassifier.

classifier
outputFile

A character(1) vector, file name for outputting prediction results. The prediction output is written to the file, tab separated.

 $assign {\tt mentCutoff}$ 

A numeric(1) vector, specifying the cutoff for classifying a putative pA site into a true or false pA class. It should be any number between 0 and 1. For example, assignmentCutoff = 0.5 will assign an putative pA site with prob\_true\_pA > 0.5 to the True class (1), and any putative pA site with prob\_true\_pA < = 0.5 as False (0).

return\_sequences

A logical(1) vector, indicating whether upstream and downstream sequences should be included in the output

#### Value

A data frame including all info as described below. The upstream and downstream sequence used in assessing the putative pA site might be included when return\_sequences = TRUE.

peak\_name the name of the putative pA site (originally from the 4th field in the bed file).

prob\_fake\_pA the probability that the putative pA site is false

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prob\_true\_pA the probability that the putative pA site is true

pred\_class the predicted class of the putative pA site, based on the assignment cutoff. 0 = Falsee/oligo(dT) internally primed, 1 = True

upstream\_seq the upstream sequence of the putative pA site used in the analysis downstream\_seq the downstream sequence of the putative pA site used in the analysis.

### Author(s)

Sarah Sheppard, Haibo Liu, Jianhong Ou, Nathan Lawson, Lihua J. Zhu

#### References

Sheppard S, Lawson ND, Zhu LJ. Accurate identification of polyadenylation sites from 3' end deep sequencing using a naive Bayes classifier. Bioinformatics. 2013;29(20):2564-2571.

```
library(BSgenome.Drerio.UCSC.danRer7)
testFile <- system.file("extdata", "test.bed",</pre>
                         package = "cleanUpdTSeq")
## convert the test set to GRanges without upstream and downstream sequence
## information
peaks <- BED6WithSeq2GRangesSeq(file = testFile,</pre>
                                skip = 1L, withSeq = TRUE)
## build the feature vector for the test set without sequence information
testSet.NaiveBayes = buildFeatureVector(peaks,
                                          genome = Drerio,
                                          upstream = 40L,
                                          downstream = 30L,
                                          wordSize = 6L,
                                          alphabet = c("ACGT"),
                                          sampleType = "unknown";
                                          replaceNAdistance = 30,
                                          method = "NaiveBayes",
                                          fetchSeq = TRUE,
                                          return_sequences = TRUE)
data(data.NaiveBayes)
## sample the test data for code testing, DO NOT do this for real data
samp <- c(1:22, sample(23:4118, 50), 4119, 4120)</pre>
Ndata.NaiveBayes <- data.NaiveBayes$Negative[, samp]</pre>
Pdata.NaiveBayes <- data.NaiveBayes$Positive[, samp]</pre>
testSet.NaiveBayes@data <- testSet.NaiveBayes@data[, samp[-1]-1]</pre>
test_out <- predictTestSet(Ndata.NaiveBayes,</pre>
                            Pdata.NaiveBayes,
                            testSet.NaiveBayes,
                            outputFile = tempfile(),
                            assignmentCutoff = 0.5)
```

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