# Package 'CODEX'

October 18, 2022

Type Package
<b>Title</b> A Normalization and Copy Number Variation Detection Method for Whole Exome Sequencing
<b>Version</b> 1.28.0
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Description A normalization and copy number variation calling procedure for whole exome DNA sequencing data. CODEX relies on the availability of multiple samples processed using the same sequencing pipeline for normalization, and does not require matched controls. The normalization model in CODEX includes terms that specifically remove biases due to GC content, exon length and targeting and amplification efficiency, and latent systemic artifacts. CODEX also includes a Poisson likelihood-based recursive segmentation procedure that explicitly models the count-based exome sequencing data.
License GPL-2
<b>Depends</b> R (>= 3.2.3), Rsamtools, GenomeInfoDb, BSgenome.Hsapiens.UCSC.hg19, IRanges, Biostrings, S4Vectors
Suggests WES.1KG.WUGSC
<b>biocViews</b> ImmunoOncology, ExomeSeq, Normalization, QualityControl, CopyNumberVariation
LazyData yes
git_url https://git.bioconductor.org/packages/CODEX
git_branch RELEASE_3_15
git_last_commit c707497
git_last_commit_date 2022-04-26
Date/Publication 2022-10-18
R topics documented:
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# **Description**

CODEX is a normalization and copy number variation calling procedure for whole exome DNA sequencing data. CODEX relies on the availability of multiple samples processed using the same sequencing pipeline for normalization, and does not require matched controls. The normalization model in CODEX includes terms that specifically remove biases due to GC content, exon length and targeting and amplification efficiency, and latent systemic artifacts. CODEX also includes a Poisson likelihood-based recursive segmentation procedure that explicitly models the count-based exome sequencing data.

### **Details**

Package: CODEX
Type: Package
Version: 0.99.0
Date: 2015-01-13
License: GPL-2

CODEX takes as input the bam files/directories for whole exome sequencing datasets and bed files for exonic positions, returns raw and normalized coverage for each exon, and calls copy number variations with genotyping results.

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### Author(s)

Yuchao Jiang <yuchaoj@wharton.upenn.edu>, Nancy R. Zhang

bambedObjDemo

Demo data pre-stored for bambedObj.

# **Description**

Pre-stored bambedObj data for demonstration purposes.

# Usage

```
data(bambedObjDemo)
```

# **Details**

Pre-computed using whole exome sequencing data of 46 HapMap samples.

#### Value

bambedObj demo data (list) pre-computed.

# Author(s)

Yuchao Jiang <yuchaoj@wharton.upenn.edu>

# **Examples**

```
bamdir <- bambedObjDemo$bamdir
sampname <- bambedObjDemo$sampname
ref <- bambedObjDemo$ref
projectname <- bambedObjDemo$projectname
chr <- bambedObjDemo$chr</pre>
```

choiceofK

Determine the number of latent factors K.

# **Description**

Determines the number of latent variables K via AIC, BIC, and deviance reduction. A pdf file containing all three plots is generated.

### Usage

```
choiceofK(AIC, BIC, RSS, K, filename)
```

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# Arguments

AIC	$vector\ of\ AIC\ for\ each\ K\ returned\ from\ {\color{red} normalize}$
BIC	vector of BIC for each $\boldsymbol{K}$ returned from $\texttt{normalize}$
RSS	vector of RSS for each $\boldsymbol{K}$ returned from $\texttt{normalize}$
K	$vector\ of\ K\ returned\ from\ {\tt normalize}$
filename	Filename of the output plot of AIC and RSS

### **Details**

AIC: Akaike information criterion, used for model selection; BIC: Bayesian information criterion, used for model selection; RSS: residue sum of squares, used to plot scree plot and determine the 'elbow'.

#### Value

pdf file with three plots: AIC, BIC, and percentage of variance explained versus the number of latent factors.

# Author(s)

Yuchao Jiang <yuchao j@wharton.upenn.edu>

### See Also

```
normalize, segment
```

# **Examples**

coverageObjDemo

Demo data pre-stored for coverageObj.

# **Description**

Pre-stored coverageObj data for demonstration purposes.

### Usage

```
data(coverageObjDemo)
```

gcDemo 5

# **Details**

Pre-computed using whole exome sequencing data of 46 HapMap samples.

# Value

coverageObj demo data (list) pre-computed.

# Author(s)

Yuchao Jiang <yuchaoj@wharton.upenn.edu>

# **Examples**

```
Y <- coverageObjDemo$Y readlength <- coverageObjDemo$readlength
```

gcDemo

Demo data pre-stored for GC content.

# Description

Pre-stored GC content data for demonstration purposes.

# Usage

```
data(gcDemo)
```

### **Details**

Pre-computed using whole exome sequencing data of 46 HapMap samples.

### Value

```
gc demo data (vector) pre-computed.
```

### Author(s)

Yuchao Jiang <yuchaoj@wharton.upenn.edu>

```
head(round(gcDemo, 2))
```

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getbambed	Get bam file directories, sample names, and exonic positions

# Description

Gets bam file directories, sample names from .txt file, and exonic positions from .bed file.

### Usage

```
getbambed(bamdir,bedFile,sampname,projectname,chr)
```

### **Arguments**

bamdir Column vector. Each line specifies directory of a bam file. Should be in same

order as sample names in sampname.

bedFile Path to bed file specifying exonic targets. Is of type character.

sampname Column vector. Each line specifies name of a sample corresponding to the bam

file. Should be in same order as bam directories in bamdir.

projectname String specifying the name of the project. Data will be saved using this as prefix.

chr Chromosome.

# Value

bamdir Bam directories sampname Sample names

ref IRanges object specifying exonic positions projectname String specifying the name of the project.

chr Chromosome

### Author(s)

Yuchao Jiang <yuchaoj@wharton.upenn.edu>

# References

Lawrence M, Huber W, Pages H, Aboyoun P, Carlson M, Gentleman R, Morgan M and Carey V (2013). "Software for Computing and Annotating Genomic Ranges." PLoS Computational Biology, 9.

#### See Also

getcoverage

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### **Examples**

```
library(WES.1KG.WUGSC)
dirPath <- system.file("extdata", package = "WES.1KG.WUGSC")
bamFile <- list.files(dirPath, pattern = '*.bam$')
bamdir <- file.path(dirPath, bamFile)
sampnameFile <- file.path(dirPath, "sampname")
sampname <- as.matrix(read.table(sampnameFile))
chr <- 22
bambedObj <- getbambed(bamdir = bamdir, bedFile = file.path(dirPath, "chr22_400_to_500.bed"), sampname = sampname, projectname = "CODEX_demo", chr)
bamdir <- bambedObj$bamdir
sampname <- bambedObj$sampname
ref <- bambedObj$ref
projectname <- bambedObj$projectname
chr <- bambedObj$chr</pre>
```

getcoverage

Get depth of coverage from whole exome sequencing

# Description

Gets depth of coverage for each exon across all samples from whole exome sequencing files.

### Usage

```
getcoverage(bambedObj, mapqthres)
```

### **Arguments**

bambedObj Object returned from getbambed

mapqthres Mapping quality threshold hold of reads.

### Value

Y Read depth matrix

readlength Vector of read length for each sample

# Author(s)

Yuchao Jiang <yuchao j@wharton.upenn.edu>

#### See Also

getbambed

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### **Examples**

```
library(WES.1KG.WUGSC)
dirPath <- system.file("extdata", package = "WES.1KG.WUGSC")</pre>
bamFile <- list.files(dirPath, pattern = '*.bam$')</pre>
bamdir <- file.path(dirPath, bamFile)</pre>
sampnameFile <- file.path(dirPath, "sampname")</pre>
sampname <- as.matrix(read.table(sampnameFile))</pre>
chr <- 22
bambedObj <- getbambed(bamdir = bamdir, bedFile = file.path(dirPath,</pre>
    "chr22_400_to_500.bed"), sampname = sampname,
    projectname = "CODEX_demo", chr)
bamdir <- bambedObj$bamdir</pre>
sampname <- bambedObj$sampname</pre>
ref <- bambedObj$ref</pre>
projectname <- bambedObj$projectname</pre>
chr <- bambedObj$chr</pre>
coverageObj <- getcoverage(bambedObj, mapqthres = 20)</pre>
Y <- coverageObj$Y
readlength <- coverageObj$readlength</pre>
```

getgc

Get GC content for each exonic target

# Description

Computes GC content for each exon. Will be later used in QC procedure and normalization.

### Usage

```
getgc(chr, ref)
```

### **Arguments**

chr Chromosome returned from getbambed
ref IRanges object returned from getbambed

#### Value

Vector of GC content for each exon.

### Author(s)

Yuchao Jiang <yuchaoj@wharton.upenn.edu>

#### References

Team TBD. BSgenome.Hsapiens.UCSC.hg19: Full genome sequences for Homo sapiens (UCSC version hg19). R package version 1.3.99.

getmapp 9

### See Also

```
getbambed, qc, normalize
```

### **Examples**

```
ref <- IRanges(st = 51207851, end = 51207982)
gc <- getgc(chr = 22, ref)</pre>
```

getmapp

Get mappability for each exonic target

# **Description**

Computes mappability for each exon. To save running time, take values from pre-computed results. Will be later used in QC procedure.

#### Usage

```
getmapp(chr, ref)
```

### Arguments

chr Chromosome returned from getbambed
ref IRanges object returned from getbambed

#### **Details**

To calculate the exonic mappability, we first construct consecutive reads of length 90 that are one base pair apart along the exon. The sequences are taken from the hg19 reference. We then find possible positions across the genome that the reads can map to allowing for two mismatches. We compute the mean of the probabilities that the overlapped reads map to the target places where they are generated and use this as the mappability of the exon.

# Value

Vector of mappability for each exon.

#### Author(s)

Yuchao Jiang <yuchao j@wharton.upenn.edu>

### See Also

```
getbambed, qc
```

```
ref <- IRanges(st = 51207851, end = 51207982)
mapp <- getmapp(chr = 22, ref)</pre>
```

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mappability

Pre-computed mappabilities

# Description

The results of pre-computed mappabilities to save running time.

# Usage

```
data(mappability)
```

# **Details**

Pre-computed mappabilities. Method used is detailed in getmapp.

# Value

List of length 24 with pre-computed mappability of each chromosome.

### Author(s)

Yuchao Jiang <yuchao j@wharton.upenn.edu>

# See Also

getmapp

# **Examples**

```
# mappability of chromosome 1
head(round(mappability[[1]], 2))
```

mappDemo

Demo data pre-stored for mappability.

# **Description**

Pre-stored mappability data for demonstration purposes.

# Usage

```
data(mappDemo)
```

# **Details**

Pre-computed using whole exome sequencing data of 46 HapMap samples.

mapp\_ref

# Value

mapp demo data (vector) pre-computed.

# Author(s)

Yuchao Jiang <yuchaoj@wharton.upenn.edu>

# **Examples**

```
head(round(mappDemo, 2))
```

mapp\_ref

Position reference for pre-computed mappability results.

# **Description**

List consisting of IRanges objects specifying exonic positions whose mappabilities are pre-computed across the genome.

### Usage

```
data(mapp_ref)
```

#### **Details**

Genomic positions for pre-computed mappabilities. Method used is detailed in getmapp.

### Value

List of length 24 with genomic positions of pre-computed mappability of each chromosome.

# Author(s)

Yuchao Jiang <yuchaoj@wharton.upenn.edu>

# See Also

getmapp

```
# mappability exon reference of chromosome 1
mapp_ref[[1]]
```

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Normalization of read depth from whole exome sequencing

# **Description**

Fits a Poisson log-linear model that normalizes the read depth data from whole exome sequencing. Includes terms that specifically remove biases due to GC content, exon capture and amplification efficiency, and latent systemic artifacts.

# Usage

```
normalize(Y_qc, gc_qc, K)
```

# **Arguments**

Y_qc	Read depth matrix after quality control procedure returned from qc
gc_qc	Vector of GC content for each exon after quality control procedure returned from
	qc
K	Number of latent Poisson factors. Can be an integer if optimal solution has
	been chosen or a vector of integers so that AIC, BIC, and RSS are computed for
	choice of optimal k.

# Value

Yhat	Normalized read depth matrix
AIC	AIC for model selection
BIC	BIC for model selection
RSS	RSS for model selection
17	N 1 C1

K Number of latent Poisson factors

# Author(s)

Yuchao Jiang <yuchaoj@wharton.upenn.edu>

# See Also

```
qc, choiceofK
```

```
Y_qc <- qcObjDemo$Y_qc
gc_qc <- qcObjDemo$gc_qc
normObj <- normalize(Y_qc, gc_qc, K = 1:5)
Yhat <- normObj$Yhat
AIC <- normObj$AIC
BIC <- normObj$BIC
RSS <- normObj$RSS
K <- normObj$K</pre>
```

normalize2

normalize2	Normalization of read depth from whole exome sequencing under the case-control setting

# **Description**

Fits a Poisson log-linear model that normalizes the read depth data from whole exome sequencing. Includes terms that specifically remove biases due to GC content, exon capture and amplification efficiency, and latent systemic artifacts. If the WES is designed under case-control setting, CODEX estimates the exon-wise Poisson latent factor using only the read depths in the control cohort, and then computes the sample-wise latent factor terms for the case samples by regression.

### Usage

```
normalize2(Y_qc, gc_qc, K, normal_index)
```

# Arguments

Y_qc	Read depth matrix after quality control procedure returned from qc
gc_qc	Vector of GC content for each exon after quality control procedure returned from qc
К	Number of latent Poisson factors. Can be an integer if optimal solution has been chosen or a vector of integers so that AIC, BIC, and RSS are computed for choice of optimal k.
normal_index	Indices of control samples.

### Value

Yhat	Normalized read depth matrix
AIC	AIC for model selection
BIC	BIC for model selection
RSS	RSS for model selection
K	Number of latent Poisson factors

# Author(s)

Yuchao Jiang <yuchaoj@wharton.upenn.edu>

### See Also

qc, choiceofK

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### **Examples**

```
Y_qc <- qcObjDemo$Y_qc
gc_qc <- qcObjDemo$gc_qc
normObj <- normalize2(Y_qc, gc_qc, K = 1:5, normal_index = seq(1, 45, 2))
Yhat <- normObj$Yhat
AIC <- normObj$AIC
BIC <- normObj$BIC
RSS <- normObj$RSS
K <- normObj$K</pre>
```

normObjDemo

Demo data pre-stored for normObj.

# Description

Pre-stored normObj data for demonstration purposes.

# Usage

```
data(normObjDemo)
```

# **Details**

Pre-computed using whole exome sequencing data of 46 HapMap samples.

### Value

normObj demo data (list) pre-computed.

### Author(s)

Yuchao Jiang <yuchao j@wharton.upenn.edu>

```
Yhat <- normObjDemo$Yhat
AIC <- normObjDemo$AIC
BIC <- normObjDemo$BIC
RSS <- normObjDemo$RSS
K <- normObjDemo$K</pre>
```

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qc

Quality control procedure for depth of coverage

#### **Description**

Applies a quality control procedure to the depth of coverage matrix both sample-wise and exon-wise before normalization.

### Usage

```
qc(Y, sampname, chr, ref, mapp, gc,cov_thresh,length_thresh,mapp_thresh,
    gc_thresh)
```

### **Arguments**

Υ Original read depth matrix returned from getcoverage Vector of sample names returned from getbambed sampname chr Chromosome. ref IRanges object specifying exonic positions returned from getbambed Vector of mappability for each exon returned from getmapp mapp Vector of GC content for each exon returned from getgc gc cov\_thresh Vector specifying the upper and lower bound of exonic median coverage threshold for QC. 20-4000 recommended. length\_thresh Vector specifying the upper and lower bound of exonic length threshold for QC. 20-2000 recommended. Scalar variable specifying exonic mappability threshold for QC. 0.9 recommapp\_thresh mended. gc\_thresh Vector specifying the upper and lower bound of exonic GC content threshold for

# **Details**

It is suggested that analysis by CODEX be carried out in a batch-wise fashion if multiple batches exist. CODEX further filters out exons that: have extremely low coverage-median read depth across all samples less than 20 or greater than 4000; are extremely short-less than 20 bp; are extremely hard to map- mappability less than 0.9; have extreme GC content-less than 20 or greater than 80. The above filtering thresholds are recommended and can be user-defined to be adapted to different sequencing protocols.

#### Value

Y\_qc Updated Y after QC

sampname\_qc Updated sampname after QC

QC. 20-80 recommended.

gc\_qc Updated gc after QC

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```
mapp_qc Updated mapp after QC
ref_qc Updated ref after QC
qcmat Matrix specifying results of exon-wise QC procedures
```

### Author(s)

Yuchao Jiang <yuchaoj@wharton.upenn.edu>

#### See Also

```
getbambed, getgc, getmapp
```

### **Examples**

```
Y <- coverageObjDemo$Y
sampname <- bambedObjDemo$sampname</pre>
chr <- bambedObjDemo$chr</pre>
ref <- bambedObjDemo$ref</pre>
gc <- gcDemo
mapp <- mappDemo</pre>
cov_thresh <- c(20, 4000)
length_thresh <- c(20, 2000)</pre>
mapp_thresh <- 0.9</pre>
gc_thresh <- c(20, 80)
qcObj <- qc(Y, sampname, chr, ref, mapp, gc, cov_thresh, length_thresh,</pre>
    mapp_thresh, gc_thresh)
Y_qc \leftarrow qc0bj\$Y_qc
sampname_qc <- qc0bj$sampname_qc</pre>
gc_qc <- qc0bj$gc_qc</pre>
mapp_qc <- qc0bj$mapp_qc</pre>
ref_qc <- qc0bj$ref_qc</pre>
qcmat <- qc0bj$qcmat</pre>
```

qcObjDemo

Demo data pre-stored for qcObj.

### **Description**

Pre-stored qcObj data for demonstration purposes.

# Usage

```
data(qcObjDemo)
```

### **Details**

Pre-computed using whole exome sequencing data of 46 HapMap samples.

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# Value

qcObj demo data (list) pre-computed.

#### Author(s)

Yuchao Jiang <yuchaoj@wharton.upenn.edu>

# **Examples**

```
Y_qc <- qcObjDemo$Y_qc
sampname_qc <- qcObjDemo$sampname_qc
gc_qc <- qcObjDemo$gc_qc
mapp_qc <- qcObjDemo$mapp_qc
ref_qc <- qcObjDemo$ref_qc
```

segment

Recursive segmentation algorithm for CNV detection and genotyping

# Description

Recursive segmentation algorithm for CNV detection and genotyping, using normalized read depth from whole exome sequencing.

# Usage

```
segment(Y_qc, Yhat, optK, K, sampname_qc, ref_qc, chr, lmax, mode)
```

# **Arguments**

Y_qc	Raw read depth matrix after quality control procedure returned from qc
Yhat	Normalized read depth matrix returned from normalize
optK	Optimal value K returned from choiceofK
K	Number of latent Poisson factors. Can be an integer if optimal solution has been chosen or a vector of integers so that AIC, BIC, and RSS are computed for choice of optimal k.
sampname_qc	Vector of sample names after quality control procedure returned from qc
ref_qc	IRanges object of genomic positions of each exon after quality control procedure returned from qc
chr	Chromosome number returned from getbambed
lmax	Maximum CNV length in number of exons returned.
mode	Can be either "integer" or "fraction", which respectively correspond to format of the returned copy numbers.

# Value

Final callset of CNVs with genotyping results.

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# Author(s)

Yuchao Jiang <yuchaoj@wharton.upenn.edu>

### See Also

normalize, choiceofK

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