

# Package ‘ExomeDepth’

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magrittr

**Suggests** knitr

**VignetteBuilder** knitr

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**Description** Calls copy number variants (CNVs) from targeted sequence data, typically exome sequencing experiments designed to identify the genetic basis of Mendelian disorders.

**License** GPL-3

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## R topics documented:

AnnotateExtra,ExomeDepth-method . . . . .	2
CallCNVs,ExomeDepth-method . . . . .	3
Conrad.hg19.common.CNVs . . . . .	4
count.everted.reads . . . . .	4
countBam.everted . . . . .	5
countBamInGRanges.exomeDepth . . . . .	6
C_hmm . . . . .	7

ExomeCount . . . . .	7
ExomeDepth-class . . . . .	8
exons.hg19 . . . . .	8
exons.hg19.X . . . . .	9
genes.hg19 . . . . .	9
get.power.betabinom . . . . .	10
getBamCounts . . . . .	10
get_loglike_matrix . . . . .	12
initialize, ExomeDepth-method . . . . .	12
plot-methods . . . . .	13
qbetabinom . . . . .	14
qbetabinom.ab . . . . .	14
select.reference.set . . . . .	15
somatic.CNV.call . . . . .	16
TestCNV, ExomeDepth-method . . . . .	17
viterbi.hmm . . . . .	17

## Index 19

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AnnotateExtra, ExomeDepth-method  
*AnnotateExtra*

---

### Description

Add annotations to a ExomeDepth object.

### Usage

```
## S4 method for signature 'ExomeDepth'
AnnotateExtra(x, reference.annotation,
  min.overlap = 0.5, column.name)
```

### Arguments

x	An ExomeDepth object.
reference.annotation	The list of reference annotations in GRanges format.
min.overlap	Numeric, defaults to 0.5. This defines the minimum fraction of the CNV call that is covered by the reference call to declare that there is a significant overlap.
column.name	The name of the column used to store the overlap (in the slot CNV.calls).

### Details

This function takes annotations in the GRanges format and adds these to the CNV calls in the ExomeDepth object. Note that a recent version of GenomicRanges (> 1.8.10) is required. Otherwise the function will return a warning and not update the ExomeDepth object.

**Value**

An ExomeDepth object with the relevant annotations added to the CNVcalls slot.

---

CallCNVs,ExomeDepth-method

*CallCNVs*

---

**Description**

Call CNV data from an ExomeDepth object.

**Usage**

```
## S4 method for signature 'ExomeDepth'
CallCNVs(x, chromosome, start, end, name,
         transition.probability = 1e-04, expected.CNV.length = 50000)
```

**Arguments**

x	An ExomeDepth object
chromosome	Chromosome information for each exon (factor).
start	Start (physical position) of each exon (numeric, must have the same length as the chromosome argument).
end	End (physical position) of each exon (numeric, must have the same length as the chromosome argument).
name	Name of each exon (character or factor).
transition.probability	Transition probability of the hidden Markov Chain from the normal copy number state to either a deletion or a duplication. The default (0.0001) expect approximately 20 CNVs genome-wide.
expected.CNV.length	The expectation for the length of a CNV. This value factors into the Viterbi algorithm that is used to compute the transition from one state to the next, which depends on the distance between exons.

**Details**

The function must be called on an ExomeDepth object. Likelihood data must have been pre-computed which should have been done by default when the ExomeDepth object was created.

This function fits a hidden Markov model to the read depth data with three hidden states (normal, deletion, duplication).

**Value**

The same ExomeDepth object provided as input but with the slot CNVcalls containing a data frame with the output of the calling.

---

Conrad.hg19.common.CNVs

*Conrad et al common CNVs*

---

### Description

Positions of common CNV calls (detected in a panel of 42 sample) from the Conrad et al paper (Nature 2010). This is build hg19 of the human genome.

### Format

A data frame with common CNV calls.

### Source

Conrad et al, Origins and functional impact of copy number variation in the human genome, Nature 2010

---

count.everted.reads     *Count the number of everted reads for a set of BAM files.*

---

### Description

This is the ExomeDepth high level function that takes a GenomicRanges object, a list of indexed/sorted BAM files, and compute the number of everted reads in each of the defined bins.

### Usage

```
count.everted.reads(bed.frame = NULL, bed.file = NULL, bam.files,
  index.files = bam.files, min.mapq = 20, include.chr = FALSE)
```

### Arguments

bed.frame	data.frame containing the definition of the regions. The first three columns must be chromosome, start, end.
bed.file	character file name. Target BED file with the definition of the regions. This file will only be used if no bed.frame argument is provided. No headers are assumed so remove them if they exist. Either a bed.file or a bed.frame must be provided for this function to run.
bam.files	character, list of BAM files to extract read count data from.
index.files	Optional character argument with the list of indexes for the BAM files, without the '.bai' suffix. If the indexes are simply obtained by adding .bai to the BAM files, this argument does not need to be specified.
min.mapq	numeric, minimum mapping quality to include a read.
include.chr	logical, if set to TRUE, this function will add the string 'chr' to the chromosome names of the target BED file.

## Details

Everted reads are characteristic of the presence of duplications in a BAM files. This routine will parse a BAM files and the suggested use is to provide relatively large bins (for example gene based, and ExomeDepth has a genes.hg19 object that is appropriate for this) to flag the genes that contain such reads suggestive of a duplication. A manual check of the data using IGV is recommended to confirm that these reads are all located in the same DNA region, which would confirm the presence of a copy number variant.

## Value

A data frame that contains the region and the number of identified reads in each bin.

## Note

This function calls a lower level function called XXX that works on each single BAM file.

## References

Computational methods for discovering structural variation with next-generation sequencing, Medvedev P, Stanciu M, Brudno M., Nature Methods 2009

## See Also

getBAMCounts

## Examples

```
## Not run:  test <- count.everted.reads (bed.frame = genes.hg19,
  bed.file = NULL,
  bam.files = bam.files,
  min.mapq = 20,
  include.chr = FALSE)

## End(Not run)
```

---

countBam.everted	<i>Counts everted reads from a single BAM file</i>
------------------	--

---

## Description

This is a utility function that is called by the higher level count.everted.reads. It processes each BAM file individually to generate the count data.

## Usage

```
countBam.everted(bam.file, granges, index = bam.file, min.mapq = 1)
```

**Arguments**

bam.file	BAM file that needs to be parsed
granges	Genomic Ranges object with the location of the bins for which we want to count the everted reads.
index	Index for the BAM files.
min.mapq	Minimum mapping quality to include reads.

**Details**

Most users will not use this function, and it will only be called by the higher level `count.everted.reads`. Nevertheless it may be useful on its own in some cases.

**Value**

A list with the number of reads in each bin.

**See Also**

`count.everted.reads`

---

`countBamInGRanges.exomeDepth`

*Compute read count data from BAM files.*

---

**Description**

Parses a BAM file and count reads that are located within a target region defined by a `GenomicRanges` object.

**Usage**

```
countBamInGRanges.exomeDepth(bam.file, index = bam.file, granges,
  min.mapq = 1, read.width = 1)
```

**Arguments**

bam.file	BAM file to be parsed
index	Index of the BAM file, without the '.bai' suffix.
granges	Genomic ranges object defining the bins
min.mapq	Minimum read mapping quality (Phred scaled).
read.width	For single end reads, an estimate of the fragment size. For paired reads, the fragment size can be directly computed from the paired alignment and this value is ignored.

**Details**

Largely derived from its equivalent function in the exomeCopy package.

**Value**

A GRanges object with count data.

---

C\_hmm

*C\_hmm*

---

**Description**

Implements the hidden Markov Model using a C routine

---

ExomeCount

*Example dataset for ExomeDepth*

---

**Description**

An example dataset of 4 exome samples, chromosome 1 only.

**Format**

A data frame with 25592 observations on the following 9 variables:

- chromosome, Character vector with chromosome names (only chromosome 1 in that case)
- start, start of exons
- end, end of exons
- exons, character name of exons
- camfid.032KA\_sorted\_unique.bam
- camfid.033ahw\_sorted\_unique.bam
- camfid.034pc\_sorted\_unique.bam
- camfid.035if\_sorted\_unique.bam
- GC, a numeric vector with the GC content

**Source**

Dataset generated in collaboration with Sergey Nejentsev, University of Cambridge.

---

ExomeDepth-class      *Class ExomeDepth*

---

### Description

A class to hold the read count data that is used by ExomeDepth to call CNVs.

### Objects from the Class

Objects can be created by calls of the form `new("ExomeDepth", data = NULL, test, reference, formula = 'cbind(test, reference) ~ 1', subset.for.speed = NULL)`. `data` is optional and is only used if the `formula` argument refers to covariates (in which case these covariates must be included in the data frame). `test` and `reference` refer to the read count data for the test and reference samples. Creating a `ExomeDepth` object will automatically fit the beta-binomial model (using routines from the `aod` package) and compute the likelihood for the three copy number states (normal, deletion and duplication).

### References

A robust model for read count data in exome sequencing experiments and implications for copy number variant calling, Plagnol et al 2012

### See Also

`?select.reference.set` `?CallCNVs`

### Examples

```
showClass("ExomeDepth")
```

---

exons.hg19      *Positions of exons on build hg19 of the human genome*

---

### Description

Exon position extracted from the ensembl database version 71.

### Format

A data frame with 192,379 observations on the following 4 variables:

- chromosome, a factor with levels 1, 2 3 4, 5 6 7 8 9, 10 11 12 13 14 15 16 17 18 19 2 20 21 22
- start a numeric vector
- end a numeric vector
- name A character vector of names for the exon(s)

**Source**

Ensemble database version 71.

---

exons.hg19.X	<i>Positions of exons on build hg19 of the human genome and on chromosome X</i>
--------------	---

---

**Description**

Exon position extracted from the ensembl database version 61 and on chromosome X only.

**Format**

A data frame of exons with the following 4 variables:

- chromosome, a factor with levels X, Y.
- start Numeric.
- end Numeric.
- name Character names for the exons.

**Source**

Ensemble database version 71.

---

genes.hg19	<i>Positions of genes on build hg19 of the human genome</i>
------------	---

---

**Description**

Exon position extracted from the ensembl database version 71.

**Format**

A data frame with 18,033 observations on the following 4 variables:

- chromosome, a factor with levels 1, 2 3 4, 5 6 7 8 9, 10 11 12 13 14 15 16 17 18 19 2 20 21 22
- start a numeric vector
- end a numeric vector
- name A character vector of names for the exon(s)

**Source**

Ensemble database version 71.

---

`get.power.betabinom`     *Estimate the power to compare two beta-binomial distributions.*

---

### Description

A power study useful in the context of ExomeDepth.

### Usage

```
get.power.betabinom(size, my.phi, my.p, my.alt.p, theory = FALSE,
  frequentist = FALSE, limit = FALSE)
```

### Arguments

<code>size</code>	Number of samples from the beta-binomial distribution.
<code>my.phi</code>	Over-dispersion parameter.
<code>my.p</code>	Expected $p$ under the null.
<code>my.alt.p</code>	Expected $p$ under the alternative.
<code>theory</code>	logical, should a theoretical limit (large sample size) be used? Defaults to FALSE.
<code>frequentist</code>	logical, should a frequentist version be used? Defaults to FALSE.
<code>limit</code>	logical, should another large sample size limit be used? Defaults to FALSE.

### Value

An expected Bayes factor.

---

`getBamCounts`     *Get count data for multiple exomes*

---

### Description

Essentially a wrapper for the accessory function `countBamInGRanges` which only considers a single BAM file at a time.

### Usage

```
getBamCounts(bed.frame = NULL, bed.file = NULL, bam.files,
  index.files = bam.files, min.mapq = 20, read.width = 300,
  include.chr = FALSE, referenceFasta = NULL)
```

**Arguments**

bed.frame	data.frame containing the definition of the regions. The first three columns must be chromosome, start, end.
bed.file	character file name. Target BED file with the definition of the regions. This file will only be used if no bed.frame argument is provided. No headers are assumed so remove them if they exist. Either a bed.file or a bed.frame must be provided for this function to run.
bam.files	character, list of BAM files to extract read count data from.
index.files	Optional character argument with the list of indexes for the BAM files, without the '.bai' suffix. If the indexes are simply obtained by adding .bai to the BAM files, this argument does not need to be specified.
min.mapq	numeric, minimum mapping quality to include a read.
read.width	numeric, maximum distance between the side of the target region and the middle of the paired read to include the paired read into that region.
include.chr	logical, if set to TRUE, this function will add the string 'chr' to the chromosome names of the target BED file.
referenceFasta	character, file name for the reference genome in fasta format. If available, GC content will be computed and added to the output.

**Details**

This function is largely a copy of a similar one available in the exomeCopy package.

**Value**

A GenomicRanges object that stores the read count data for the BAM files listed as argument.

**Author(s)**

Vincent Plagnol

**References**

exomeCopy R package.

**Examples**

```
## Not run:
load(exons.hg19)

my.counts <- getBamCounts(bed.frame = exonpos,
                          bam.files = my.bam,
                          referenceFasta = 'human_g1k_v37.fasta')

## End(Not run)
```

---

```
get_loglike_matrix    get_loglike_matrix
```

---

### Description

Computes the loglikelihood matrix for the three states and each exon

---

```
initialize,ExomeDepth-method  
ExomeDepth initialization tool
```

---

### Description

Builds an exomeDepth object from test and reference vectors

### Usage

```
## S4 method for signature 'ExomeDepth'  
initialize(.Object, data = NULL, test, reference,  
          formula = "cbind(test, reference) ~ 1", phi.bins = 1,  
          prop.tumor = 1, subset.for.speed = NULL, verbose = TRUE)
```

### Arguments

.Object	ExomeDepth object
data	Data frame containing potential covariates.
test	Numeric, vector of counts for the test sample.
reference	Numeric, vector of counts for the reference sample.
formula	Linear model to be used when fitting the data.
phi.bins	Numeric, defaults to 1. Number of discrete bins for the over-dispersion parameter phi, depending on read depth. Do not modify this parameter for the standard use of ExomeDepth.
prop.tumor	Numeric, defaults to 1. For the somatic variant calling, this assesses the proportion of the test sample data originating from the tumour. Do not modify this parameter for the standard use of ExomeDepth.
subset.for.speed	Numeric, defaults to NULL. If non-null, this sets the number of data points to be used for an accelerated fit of the data.
verbose	Logical, controls the output level.

plot-methods

*Plotting function for ExomeDepth objects***Description**

Plot function for the ExomeDepth class

**Usage**

```
## S4 method for signature 'ExomeDepth,ANY'
plot(x, sequence, xlim, ylim = NULL,
     count.threshold = 10, ylab = "Observed by expected read ratio",
     xlab = "", type = "b", pch = "+", with.gene = FALSE,
     col = "red", ...)
```

**Arguments**

x	ExomeDepth object
sequence	character, Name of the sequence/chromosome of the region to plot (for example "chr5" would be typical)
xlim	numeric of size 2, start and end position of the region to plot
ylim	numeric of size 2, range for the y-axis
count.threshold	numeric, minimum number of reads in the reference set to display a point in the plot
ylab	Defaults to ""
xlab	Defaults to ""
type	Defaults to 'b'
pch	Defaults to '+'
with.gene	Logical, defaults to FALSE, Should the gene information (obtained from the annotation data) be plotted under the read depth plot?
col	character, Colour for the line displaying the read depth ratio for each exon
...	Additional arguments to be passed to the base plot function

qbetabinom

*Quantile for betabin function*

---

**Description**

Quantile function for the betabinomial distribution using the  $p/\phi$  parameterisation.

**Usage**

```
qbetabinom(p, size, phi, prob)
```

**Arguments**

p	Point of the distribution from which one is looking for the quantile
size	Sample size of the random variable
phi	Over-dispersion parameter
prob	Mean probability of the binomial distribution

**Details**

Filling a gap in the VGAM package.

**Value**

A real number corresponding to the quantile  $p$ .

**Author(s)**

Vincent Plagnol

**See Also**

VGAM R package.

---

qbetabinom.ab

*Quantile function for the beta-binomial distribution*

---

**Description**

Standard qbetabinomial.ab function which is missing from the VGAM package.

**Usage**

```
qbetabinom.ab(p, size, shape1, shape2)
```

**Arguments**

p	Mean value of the beta-binomial distribution.
size	Size of the beta-binomial.
shape1	First parameter of the beta distribution for p.
shape2	Second parameter of the beta distribution for p.

**Value**

A quantile of the distribution.

**See Also**

VGAM package.

---

select.reference.set *Combine multiple samples to optimize the reference set in order to maximise the power to detect CNV.*

---

**Description**

The power to detect copy number variant (CNVs) from targeted sequence data can be maximised if the most appropriate set of sequences is used as reference. This function is designed to combine multiple reference exomes in order to build the best reference set.

**Usage**

```
select.reference.set(test.counts, reference.counts, bin.length = NULL,
  n.bins.reduced = 0, data = NULL,
  formula = "cbind(test, reference) ~ 1", phi.bins = 1)
```

**Arguments**

test.counts	Read count data for the test sample (numeric, typically a vector of integer values).
reference.counts	Matrix of read count data for a set of additional samples that can be used as a comparison point for the test sample.
bin.length	Length (in bp) of each of the regions (often exons, but not necessarily) that were used to compute the read count data (i.e. what is provided in the argument test.counts of this function). If not provided all bins are assumed to have equal length.
n.bins.reduced	This optimization function can be slow when applied genome-wide. For the purpose of building the reference sample, it is not necessary to use the full data. The number provided by this argument specifies the number of regions (typically exons) that will be sub-sampled (using a grid) to optimise the referenceset. I find that 10,000 is largely sufficient for exome data.

data	Defaults to NULL: A data frame of covariates that can be included in the model.
formula	Defaults to 'cbind(test, reference) ~ 1'. This formula will be used to fit the read count data. Covariates present in the data frame (for example GC content) can be included in the right hand side of the equation'. If covariates are provided they must be provided as arguments (in the data frame "data").
phi.bins	Numeric integer (typically 1, 2, or 3) that specifies the number of windows where the over-dispersion parameter phi can vary. It defaults to 1, i.e. a single over-dispersion parameter, independently of read depth.

**Value**

reference.choice	character: list of samples selected as optimum reference set.
summary.stats	A data frame summarizing the output of this computation, including expected Bayes factor, Rs statistic (see reference for explanation) for multiple choices of reference set.

---

somatic.CNV.call	<i>somatic.CNV.call</i>
------------------	-------------------------

---

**Description**

Call somatic variants between healthy and disease tissues.

**Usage**

```
somatic.CNV.call(normal, tumor, prop.tumor = 1, chromosome, start, end,
names)
```

**Arguments**

normal	Read count data (numeric vector) for the normal tissue.
tumor	Read count data (numeric vector) for the tumor.
prop.tumor	Proportion of the tumour DNA in the tumour sample (between 0 and 1, and less than 1 if there is normal tissue in the tumor sample).
chromosome	Chromosome information for the bins.
start	Start position of each bin (typically in bp).
end	End position of each bin.
names	Names for each bin (typically exon names but any way to track the bins will do).

**Details**

Use read depth data from targeted sequencing experiments to call CNV between a tumor and matched healthy tissue. This is an experimental function at this stage.

**Value**

An ExomeDepth object with CNV calls.

**Note**

Absolutely experimental, not the main function from the package.

---

TestCNV,ExomeDepth-method  
*TestCNV*

---

**Description**

Computes the Bayes Factor in favour of a CNV defined by position and type.

**Usage**

```
## S4 method for signature 'ExomeDepth'
TestCNV(x, chromosome, start, end, type)
```

**Arguments**

x	ExomeDepth object
chromosome	Character, chromosome name.
start	Numeric, start of the tested CNV
end	Numeric, end of the tested CNV
type	Character, must be either deletion or duplication.

---

viterbi.hmm                      *Computes the Viterbi path for a hidden markov model*

---

**Description**

Estimates the most likely path for a hidden Markov Chain using the maximum likelihood Viterbi algorithm.

**Usage**

```
viterbi.hmm(transitions, loglikelihood, positions, expected.CNV.length)
```

**Arguments**

<code>transitions</code>	Transition matrix
<code>loglikelihood</code>	numeric matrix containing the loglikelihood of the data under the possible states
<code>positions</code>	Positions of the exons
<code>expected.CNV.length</code>	Expected length of CNV calls, which has an impact on the transition matrix between CNV states.

**Details**

Standard forward-backward Viterbi algorithm using a precomputed matrix of likelihoods.

**Value**

<code>comp1</code>	Description of 'comp1'
<code>comp2</code>	Description of 'comp2'

# Index

## \*Topic **classes**

ExomeDepth-class, 8

## \*Topic **datasets**

Conrad.hg19.common.CNVs, 4

ExomeCount, 7

exons.hg19, 8

exons.hg19.X, 9

genes.hg19, 9

AnnotateExtra

(AnnotateExtra, ExomeDepth-method),  
2

AnnotateExtra, ExomeDepth-method, 2

C\_hmm, 7

CallCNVs (CallCNVs, ExomeDepth-method), 3

CallCNVs, ExomeDepth-method, 3

Conrad.hg19.common.CNVs, 4

count.everted.reads, 4

countBam.everted, 5

countBamInGRanges.exomeDepth, 6

ExomeCount, 7

ExomeDepth-class, 8

exons.hg19, 8

exons.hg19.X, 9

genes.hg19, 9

get.power.betabinom, 10

get\_loglike\_matrix, 12

getBamCounts, 10

initialize, ExomeDepth-method, 12

plot, ANY-method (plot-methods), 13

plot, ExomeDepth, ANY-method

(plot-methods), 13

plot, ExomeDepth-method (plot-methods),

13

plot-methods, 13

plot.ExomeDepth (plot-methods), 13

qbetabinom, 14

qbetabinom.ab, 14

select.reference.set, 15

somatic.CNV.call, 16

TestCNV (TestCNV, ExomeDepth-method), 17

TestCNV, ExomeDepth-method, 17

viterbi.hmm, 17