# Package 'chimera'

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<b>Depends</b> Biobase, GenomicRanges (>= 1.13.3), Rsamtools (>= 1.13.1), GenomicAlignments, methods, AnnotationDbi, BSgenome.Hsapiens.UCSC.hg19, TxDb.Hsapiens.UCSC.hg19.knownGene, Homo.sapiens
Suggests BiocParallel, geneplotter
Enhances Rsubread, BSgenome.Mmusculus.UCSC.mm9,     TxDb.Mmusculus.UCSC.mm9.knownGene,     BSgenome.Mmusculus.UCSC.mm10,     TxDb.Mmusculus.UCSC.mm10.knownGene, Mus.musculus,     BSgenome.Hsapiens.NCBI.GRCh38,     TxDb.Hsapiens.UCSC.hg38.knownGene
Description  This package facilitates the characterisation of fusion products events. It allows to import fusion data results from the following fusion finders: chimeraScan, bellerophontes, deFuse, Fusion Finder, FusionHunter, mapSplice, tophat-fusion, FusionMap, STAR, Rsubread, fusionCatcher.
biocViews Infrastructure
<b>SystemRequirements</b> STAR, TopHat, bowtie and samtools are required for some functionalities
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chimera-package

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#### **Description**

The package imports fusion results from tophat-fusion, tophat-fusion-post, mapSplice, deFuse, fusionmap, bellerophontes, fusionfinder, fusionhunter, STAR, Rsubread, fusionCatcher. The package was design to facilitate the characterisation of fusion products events. Data upload: outputs for the above indicated fusion detection tools can be imported using importFusionData in a list of fSet objects. fSet-class offers various methods to extract information from the fSet objects. The fusion names can be extracted with fusionName function. The number of reads supporting a fusion event cam be extracted with the supportingReads function.

Filtering: The imported fusion list can be filtered using filterList

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Annotation: Oncofuse can be installed in chimera with the function oncofuseInstallation Various information on the fusions location, on structural and functional domains affected by the fusion event as well as a prediction of the putative functional effect of the fusion on the cell can be obtained by using oncofuseRun.

chimeraSeqs generates the nucleotide sequence of a fusion transcript described in an fSet object. chimeraSeqSet does the same but on a list of fSet objects.

fusionPeptides allows to investigate if the fusion events generate also a fusion at protein level.

subreadRun allows to remap reads on the fused transcripts reconstructed with chimeraSeqs

Validation GapFiller is a seed-and-extend local assembler capable to produce (in-silico) longer and highly accurate sequences from a collection of Next Generation Sequencing reads. It can be installed in chimera with the function gapfillerInstallation. The function gapfillerRun allows to check if reads mapped by subreadRun over a fused transcript generated with chimeraSeqs are able to reconstruct by de novo assembly the fusion break-point.

Export The function prettyPrint converts the information stored in a list of fSet objects in a dataframe structure that is saved as tab delimited file.

#### **Details**

Package: chimera Type: Package Version: 1.0

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#### Author(s)

Raffaele A Calogero Maintainer: Raffaele A Calogero <raffaele.calogero@unito.it>

# References

Beccuti M, et al. The structure of state-of-art gene fusion-finder algorithms. OA Bioinformatics 2013;1(1):2. Carrara, M., et al. State of art fusion-finder algorithms are suitable to detect transcription-induced chimeras in normal tissues? BMC bioinformatics 2013;14 Suppl 7:S2. Carrara, M., et al. State-of-the-Art Fusion-Finder Algorithms Sensitivity and Specificity. BioMed research international 2013;2013:340620. Shugay, M., et al. Oncofuse: a computational framework for the prediction of the oncogenic potential of gene fusions. Bioinformatics 2013;29(20):2539-2546. Nadalin, F., et al. GapFiller: a de novo assembly approach to fill the gap within paired reads. BMC bioinformatics 2012;13 Suppl 14:S8.

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bam2fastq	A function to extract pair end reads from the bam file generated with subreadRun function

# **Description**

A function to extract pair end reads from the bam file generated with subread function. The output files are ready to be used for fusion validation with gapfiller

#### Usage

```
bam2fastq(bam, filename="ready4gapfiller",ref,parallel=FALSE)
```

# **Arguments**

name of the bam file to be used for PE reads extraction

filename base name for the PE fastq output data

ref name of the fusion sequence that was used as reference

parallel option that allow the use of BioParallel package

# Value

PE fastq files

# Author(s)

Raffaele A Calogero

```
#if(require(Rsubread)){
# subreadRun(ebwt=paste(find.package(package="chimera"),"/examples/SULF2_ARFGEF2.fa",sep=""),
# input1=paste(find.package(package="chimera"),"/examples/mcf7_sample_1.fq",sep=""),
# input2=paste(find.package(package="chimera"),"/examples/mcf7_sample_2.fq",sep=""),
# outfile.prefix="accepted_hits", alignment="se", cores=1)
# ref.name <- names(readDNAStringSet(paste(find.package(package="chimera"),"/examples/SULF2_ARFGEF2.fa",sep="
# bam2fastq(bam="accepted_hits.bam", filename="ready4gapfiller", ref=ref.name, parallel=F)
#}</pre>
```

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# Description

A function to extract the reads overlapping to fusion breakpoint.

# Usage

```
breakpointOverlaps(fset, plot=FALSE, ylim=NULL)
```

# **Arguments**

fset	An fSet object. The slots fusionRNA and fusionGA needs to be loaded
plot	If FALSE plot is not printed
ylim	If NULL it uses the full fusion transcript coverage to define the ylim of the plot. If setted it can be used to zoom in the plot to better see the structure of the coverage at the brek point

#### Value

An object of GAlignment class. A plot of the fusion trascript coverage in blue and of the reads spanning over the break point in yellow.

# Author(s)

Raffaele A Calogero

## **Examples**

```
load(paste(find.package="chimera"), "/examples/fset_ARFGEF2-SULF2.rda", sep=""))
my.seq <- chimeraSeqs(my.fset)
my.fset <- addRNA(my.fset, my.seq)
tmp <- breakpointOverlaps(my.fset)</pre>
```

 ${\it chimeraSeqs} \qquad \qquad {\it A function \ to \ generate \ the \ nucleotide \ sequences \ of \ a \ fusion \ event}$ 

# Description

A function generating the nucleotide sequences of a chimera.

```
chimeraSeqs(fset, extend=1000, type="transcripts")
```

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#### **Arguments**

fset A fSet object.

extend number of nucleotides used to extend a genomic region that is not an annotated

gene. Default is 1000 nts

type Chimera can be build at transcript level, i.e. transcript level will generate multi-

ple fusion transcripts depending of the number of transcripts associated to each

of the two genes involved in the fusion.

#### Value

A DNAStringSet encompassing the fusions generated using all the isoforms for each gene involved in the fusion. The name of each element of the DNAStringSet has the following format: gene1-lengthOfGeneFragment:gene2-lengthOfGeneFragment. In case the fusion junction is located in an intronic sequence, a warning is provided. The presence of a partial intron in the fusion is an indication that the fusion does not generate an active chimeric peptide.

#### Author(s)

Raffaele A Calogero

#### See Also

fusionName, chimeraSeqSet

# **Examples**

```
tmp <- importFusionData("fusionmap", paste(find.package(package="chimera"),"/examples/mcf7.FMFusionReport", sep
fusion.names <- fusionName(tmp)
fusion.names
myset <- tmp[[13]]
tmp.seq <- chimeraSeqs(myset, type="transcripts")
#writeXStringSet(tmp.seq, paste(sub(":","_",fusion.names[[13]]),".fa",sep=""), format="fasta")</pre>
```

chimeraSeqSet

A function to generates DNAStringSet encompassing fusion sequences

# Description

A function generating the nucleotide sequences of chimeras described in a list of fSet, i.e. the list generated using importFusionData function.

```
chimeraSeqSet(list, parallel=FALSE)
```

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### Arguments

list A list of fSet objects.

parallel If TRUE uses the BioParallel package

#### Value

A DNAStringSet encompassing the fusions described in a list of fSet objects. This object represents the ideal reference to remap reads over detected fusions. Remapping is required to validate fusions using GapFiller de novo reconstruction.

#### Author(s)

Raffaele A Calogero

#### See Also

```
fusionName, importFusionData, gapfillerInstallation, gapfillerRun
```

# **Examples**

```
tmp <- importFusionData("fusionmap", paste(find.package(package="chimera"),"/examples/mcf7.FMFusionReport", sep
fusion.names <- fusionName(tmp)
fusion.names
myset <- tmp[1:3]
tmp.seq <- chimeraSeqSet(myset, parallel=FALSE)
# sapply(tmp.seq, function(x){writeXStringSet(x, "detected.fusions.fa", format="fasta", append=TRUE)})</pre>
```

defuseTPTN

A function that genererate a list of fSet objects encopassing 60 experimentally validated fusions and 61 false fusions

# Description

The function uses the file defuse\_TP\_FP.txt located in the examples folder of chimera to generate a list of fSet object encompassing 60 TP fusions and 61 TN fusions extracted from supplementary table 8 of the deFuse paper McPherson et al. (2011) PLoS Comput Biol 7(5): e1001138. The fset object can be used to evaluate the efficacy of filtering approaches available in chimera package.

# Usage

```
defuseTPTN()
```

## Value

An list of fSet objects: Experimentally validated fusions 1-60, fish validated fusions 1-14, false fusions 61-121.

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#### Author(s)

Raffaele A Calogero

#### **Examples**

tptn <-defuseTPTN()</pre>

filterList

A function to filter a list of fSet objects

#### Description

A function filtering a list of fSet objects on the basis of various parameters.

### Usage

filterList(x,type=c("spanning.reads", "fusion.names", "intronic", "annotated.genes", "read.through",

## Arguments

x a list of fSet object

type filtering can be performed on the basis of spanning.read: a minimal number of

spanning reads, fusion.names: a vector list of user defined fusion names, intronic: only fusion encompassing exon data are retained annotated.genes: only fusion encompassing two annotated genes are retained read.through: only fusion with different gene names are retained oncofuse: using the output of oncofuseRun various filtering option can be applied using oncofuse.type and oncofuse.

fuse.threshold

oncofuse.output

The output generated with oncofuseRun

query is a number. In case spanning reads is selected or a vector of fusion names

if the case fusion.names is selected. In case type is intronic query is NULL. In the latter case fusion having one of the elements located in an intronic region are discarded. In the case of oncofuse.type equal to passenger.prob or expres-

sion.gain query is the threshold number to be used for the filtering

oncofuse.type This refers to the filtering based on the output of oncofuse that can be generated

using the oncofuseRun function. g5CDS selects only fusions having the breakpoint in 5' end gene CDS, g3CDS selects only fusions having the breakpoint in 3' end gene CDS, g5g3CDS selects only fusions having the breakpoint in both gene CDSs, g5exon selects only fusions having the breakpoint in an exon of 5' end gene, g3exon selects only fusions having the breakpoint in an exon of 3' end gene, g5g3exon selects only fusions having the breakpoint in an exon of both genes. In the case of oncofuse type equal to driver prob the filter will use the probability of the fusion to be a tumor driver. In the case of oncofuse type equal to expression gain the filter will use the score that suggests the presence

of a gain in expression

parallel option to run a parallel version of the function, default FALSE

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#### Author(s)

Raffaele A Calogero

#### **Examples**

```
tmp <- importFusionData("fusionmap", paste(find.package(package="chimera"),"/examples/mcf7.FMFusionReport", sep
fusion.names <- fusionName(tmp)
tmp1 <- filterList(tmp, type="fusion.names", fusion.names[c(1,3,7)], parallel=FALSE)
tmp2 <- filterList(tmp, type="spanning.reads", query=2, parallel=FALSE)
#tmp3 <- filterList(tmp, type="intronic")
#tmp4 <- filterList(tmp, type="annotated.genes", parallel=FALSE)
#tmp5 <- filterList(tmp, type="read.through", parallel=FALSE)
#csdf.of <- oncofuseRun(csdf.e, tissue="EPI")
#tmp6 <- filterList(csdf.e[1:100],oncofuse.output=csdf.of, type="oncofuse", oncofuse.type="g5g3CDS", parallel=T
#tmp7 <- filterList(csdf.e[1:100],oncofuse.output=csdf.of, type="oncofuse", oncofuse.type="passenger.prob", que</pre>
```

filterSamReads

A function to filter SAM or BAM files

## **Description**

A function to filter SAM or BAM files using picard-tools

#### Usage

```
filterSamReads(input, output, filter=c("includeAligned","excludeAligned"))
```

#### **Arguments**

input SAM/BAM file to be validated

output file name in which to save the filtered reselts

filter type of filter

#### Value

A filtered SAM/BAM.

#### Author(s)

Raffaele A Calogero

#### See Also

picardInstallation

### **Examples**

# filterSamReads(input="kd2\_accepted\_hits2.sam", output="kd2\_accepted\_hits2\_mapped.sam", filter="includeAlignec

10 fSet

fSet Class fSet, a class represent fusion data, and methods for processing it

# Description

This is class representation for a fusion event.

#### **Slots**

fusionInfo A list: fusionTool: the tool that has generated the fusions UniqueCuttingPositionCount: the number of unique cutting positions detected for the fusion. SeedCount: the number of reads overlapping the break-point, i.e. spanning reads (FusionMap, FusionHunter, mapSplice, Tophat-fusion, ChimeraScan, STAR, Rsubread, FusionCatcher). Both spanning and encompassing reads (Bellerophontes, FusionFinder). Encompassing reads, i.e. one read of a pair on gene 1, and the other on gene2 (deFuse). RescuedCount: the number of reads overlapping the break-point rescued after identification of the break point (FusionMap). Encompassing reads (Tophat-fusionm Tophat-fusion-post, FusionCatcher, STAR). Both spanning and encompassing reads (ChimeraScan, Rsubread). SplicePattern: the splice pattern for a fusion junction FusionGene: the name of the fusion gene in the format gene1 -> gene2. frameShift: frameshift at break-point

fusionLoc A GRangesList encompassing fusion locations for gene 1 and 2

**fusionRNA** A DNAStringSet encompassing the fusion transcript that can be generated by chimeraSeqs function.

**fusionGA** A GAlignments object encompassing positions for all reads mapping on the DNAS-tringSet located in fusionRNA slot

#### Methods

Standard generic methods:

fusionData(fSet) An accessor function used to retrieve information for a fusion

**fusionGRL(fSet)** An accessor function used to retrieve GRangesList encompassing fusion locations for gene 1 and 2

fusionRNA(fSet) An accessor function used to extract the DNAStringSet

addRNA(fSet, rna) An accessor function used to add the DNAStringSet to the fset object

**fusionGA(fSet)** An accessor function used to extract the GAlignments object

addGA(fSet, bam) An accessor function used to add the GAlignments object to the fSet object

#### Constructor

fusionInfo A list of the fusin characteristics see above slot fusionInfo fusionLoc A GRangesList encompassing the fusion break points

fusionRN

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fusionRNA A DNAStringSet encompassing the fusion transcript fusionGA A GAlignments encompassing the location of reads mapping on the fusion transcript

# Author(s)

Raffaele A Calogero

#### See Also

chimeraSegs, oncofuseRun, fusionPeptides, prettyPrint

#### **Examples**

```
#creating a fusion report from output of fusionMap
tmp <- importFusionData("fusionmap", paste(find.package(package="chimera"),"/examples/mcf7.FMFusionReport", sep</pre>
#extracting the fusions names
fusion.names <- fusionName(tmp)</pre>
fusion.names
#extracting the fSet object ofr one of the fusions
myset \leftarrow tmp[[13]]
#constructing the fused sequence(s)
trs <- chimeraSeqs(myset, type="transcripts")</pre>
#adding the sequences to the fSet object
myset <- addRNA(myset , trs)</pre>
#extracting sequences from an fSet object
tmp.seq <- fusionRNA(myset)</pre>
#adding reads mapped on the fusion generated using tophatRun function
myset <- addGA(myset, paste(path.package(package="chimera"),"/examples/mcf7_trs_accepted_hits.bam",sep=""))</pre>
#extracting the GAlignments from an fSet object
ga <- fusionGA(myset)</pre>
```

fusionName

A function to extract fusion names for a list of fSet object

# **Description**

A function allowing extract fusion names from a list of fSet objects.

# Usage

```
fusionName(list, parallel=FALSE)
```

# Arguments

list a list of fSet object

parallel option to run a parallel version of the function, default FALSE

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#### Author(s)

Raffaele A Calogero

#### **Examples**

```
tmp <- importFusionData("fusionmap", paste(find.package(package="chimera"),"/examples/mcf7.FMFusionReport", sep
fusion.names <- fusionName(tmp)
fusion.names</pre>
```

fusionPeptides

A function to investigate the peptides involved in the fusion event.

### **Description**

A function extracting the donor and the acceptor peptides involved in the fusion.

#### Usage

fusionPeptides(chimeraSeq.output, annotation="hsUCSC")

#### **Arguments**

chimeraSeq.output

DNAStringSet encompassing the fusion event of interest, generated by chimeraSeq

function

annotation The annotation used to retrieve the UCSC names of the transcripts involved in

the fusion

#### Value

An list encompassing:

AAStringSet encompassing: fusion sequence, peptide from p1 and peptide from p2. In case

the peptides are not in frame the AAStringSet will not contain the fusion se-

quence

tmpx <- lapply(tmp.seq,fusionPeptides)</pre>

DNAStringSet encompassing the fusion transcript

#### Author(s)

Raffaele A Calogero

```
tmp <- importFusionData("fusionmap", paste(find.package(package="chimera"),"/examples/mcf7.FMFusionReport", sep
fusion.names <- fusionName(tmp)
fusion.names
myset <- tmp[1:3]
tmp.seq <- chimeraSeqSet(myset, parallel=FALSE)</pre>
```

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gapfillerInstallation A function to download a compiled version of GapFiller

# Description

A function allowing the download and installation of a compiled version of GapFiller in chimera package folder. The source code of GapFiller can be retrieved at http://sourceforge.net/projects/gapfiller/files/?source=navbar. The paper describing the tool is Nadalin F, Vezzi F, Policriti A. GapFiller: a de novo assembly approach to fill the gap within paired reads. BMC Bioinformatics. 2012;13 Suppl 14:S8.

# Usage

```
gapfillerInstallation(os=c("mac64", "unix64"))
```

#### **Arguments**

os

The supported operating systems

#### Author(s)

Raffaele A Calogero

#### See Also

gapfillerRun

# **Examples**

```
#gapfillerInstallation(os="mac64")
```

gapfillerRun

A function to confirm fusion break point by de novo assembly

# Description

A function that uses GapFiller to confirm, by de novo assembly, the presence of the fusion break point. The function needs as input the fusion transcript generated by chimeraSeqs function and two fastq files corresponding to the reads mapping over the fusion transcript.

```
gapfillerRun(fusion.fa, seed1, seed2, gapfiller=NULL, seed.ins=200, seed.var=50,
block.length=5, overlap=20, max.length=5000, slack=30,
k=6, global.mismatch=5)
```

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### **Arguments**

fusion. fa fasta file with the fusin transcript

seed1 The R1 fastq of a pair-end seed2 The R2 fastq of a pair-end

gapfiller path to GapFiller executable program

seed.ins seed reads insert size

seed.var seed reads insert variation block.length length of perfect match

overlap minimum suffix-prefix overlap

max.length print only contigs <= max-length long

slack number of overlaps: suffix-prefix overlap range is given by overlap, overlap +

slack

k length of the word used to hash

global.mismatch

maximum number of mismatches between mate and contig

#### Value

The program will write in a temporary directory contigs.fasta and contig.stats, which are used to evaluate if the de novo assembly allows the identification of the fusion break point. The function returns a list of three objects. The list is returned only in case that some of de novo assemblies cover the breakpoint junction. The list is made of:

contigs which is a Pairwise Alignments object

junction.contigs

which is a DNAStringSet encompassing the sequences present in the contigs

object

fusion which is a DNAStringSet object encompassing the fusion transcript

# Author(s)

Raffaele A Calogero

#### See Also

chimeraSeqs, gapfillerInstallation

#### **Examples**

# tmp <- gapfillerRun(fusion.fa=paste(path.package("chimera", quiet = FALSE),"/examples/uc002xvp.1-243\_uc002iyu

gapfillerWrap 15

#### **Description**

A function that uses GapFiller to confirm, by de novo assembly, the presence of the fusion break point. The function needs as input a list of fusion transcript generated by chimeraSeqSet function and the bam file containing the reads remapped over the fusion transcripts made using subreadRun.

# Usage

```
gapfillerWrap(chimeraSeqSet.out, bam, parallel=c(FALSE,TRUE))
```

# **Arguments**

chimeraSeqSet.out

a list of DNAStringSet output from chimeraSeqSet

bam file containing the reads remapped over the fusion transcripts using Rsub-

read

parallel if FALSE FALSE no parallelization, if TRUE TRUE full parallelization, if FALSE

TRUE only parallelization for internal funtions

#### Value

The program will write in a temporary directory contigs.fasta and contig.stats, which are used to evaluate if the de novo assembly allows the identification of the fusion break point. The function returns for each fusion a list of three objects. The list is returned only in case that some of de novo assemblies cover the breakpoint junction. The list is made of:

```
contigs which is a PairwiseAlignments object junction.contigs
```

which is a DNAStringSet encompassing the sequences present in the contigs

object

fusion which is a DNAStringSet object encompassing the fusion transcript

# Author(s)

Raffaele A Calogero

# See Also

```
chimeraSeqs, gapfillerInstallation, gapfillerRun
```

```
#tmp <- importFusionData("star", "Chimeric.out.junction", org="hg19", min.support=100)
#myset <- tmp[1:4]
#tmp.seq <- chimeraSeqsSet(myset, type="transcripts")
#tmp <- gapfillerWrap(chimeraSeqSet.out=trsx, bam="accepted_hits_mapped.bam", parallel=c(FALSE,TRUE))</pre>
```

16 importFusionData

importFusionData

A function to import fusion data detected by different fusion finders

#### **Description**

A function to import in a list fusions data detected by bellerophontes, defuse, fusionfinder, fusion-hunter, mapsplice, tophat-fusion, fusionmap, chimerascan, STAR, Rsubread, fusionCatcher. In the case of chimerascan and STAR output it is possible to load the data applying a filter on the minimal number of reads supporting a specific fusion. Both chimerascan and STAR accept data generated using human hg19, hg38, mouse mm9 and mm10 reference. IMPORTANT: please note that the it is important that the genome reference version used for the alignment is the same of that used for by chimera for annotation. Expecially between hg38 and hg19 there are shifts in gene location.

#### Usage

```
importFusionData(format, filename, ...)
```

#### **Arguments**

format

Format allows to select the data structure to be imported. One of the following keyword need to be associated to format parameter: bellerophontes, defuse, fusionfinder, fusionhunter, mapsplice, tophat-fusion, fusionmap, chimerascan, star, rsubread, fusioncatcher.

filename

The file generated by one of the fusion finders defined in format

. . .

Additional parameters: In case of rsubread min.distance, which indicates the minimal distance to consider a fusion within the same chromosome, is set to 700000 as default. In case of rsubread, chimerascan and STAR min.support, which indicates the minimal number of reads supporting a specific fusion, is set to 10 as default. Please remember that using low values as min.support, e.g. 1, will significantly increase the time for data import. In this case is strongly suggested to run the function as batch and, when available, using the parallel option, see below. In case of chimerascan, STAR, Rsubread, fusionmap, fusioncatcher, tophat-fusion, tophat-fusion-post, mapsplice, bellerophontes org, which indicates the organism to be used for annotation, has the following options: hg19, hg38, mm9, mm10. In case of STAR and Rsubread the upload can be parallelized using parallel=TRUE

#### Author(s)

Raffaele A Calogero

# **Examples**

tmp <- importFusionData("fusionmap", paste(find.package(package="chimera"),"/examples/mcf7.FMFusionReport",
#min.support allow to retrieve only the subset of fusions supported by a user defined minimal number of junction
#tmp <- importFusionData("chimerascan", "edgren\_cs10.bedpe", min.support=10, org="hg19")
#download Edgren Chimeric.out.junction. This file encompass the results obtaines combined all cell lines used</pre>

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```
#download.file("http://sourceforge.net/projects/ochguiextras/files/chimera/Chimeric.out.junction.zip/download.file("http://sourceforge.net/projects/ochguiextras/files/chimera/Chimeric.out.junction.zip/download.graphy.com/chimeric.out.junction.zip")
#tmp <- importFusionData("star", "Chimeric.out.junction", org="hg19", min.support=100)</pre>
```

is.fSet

A function to evaulate if an object belongs to fSet class or not

# Description

A function to evaulate if an object belongs to fSet class or not.

# Usage

```
is.fSet(x)
```

#### **Arguments**

Х

an object

#### Value

If the object belons to fSet class it returns TRUE, else it returned FALSE

# Author(s)

Raffaele A Calogero

# **Examples**

```
tmp <- importFusionData("fusionmap", paste(find.package(package="chimera"),"/examples/mcf7.FMFusionReport", sep
is.fSet(tmp[[1]])
```

MHmakeRandomString

A function generating a random string

# Description

A function generating a random string.

# Usage

MHmakeRandomString()

#### Value

a string

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#### Author(s)

Raffaele A Calogero

#### **Examples**

```
tmp.file <- paste(MHmakeRandomString(),".fa", sep="")</pre>
```

newfSet

A constructor for fSet class objects

# **Description**

A function to create a new fSet object

#### Usage

# **Arguments**

fusionInfo A list of the fusion characteristics see fSet class slot fusionInfo

fusionLoc A GRangesList encompassing the fusion break points fusionRNA A DNAStringSet encompassing the fusion transcript

fusionGA A GAlignments encompassing the location of reads mapping on the fusion tran-

script

#### Value

An object of fSet class

#### Author(s)

Raffaele A Calogero

```
tmp <- newfSet()
tmp</pre>
```

oncofuseInstallation 19

 $oncofuseInstallation \ A \ function \ to \ download \ oncofuse$ 

# Description

A function allowing the download of oncofuse in the chimera folder. Oncofuse requires java

# Usage

```
oncofuseInstallation()
```

# Author(s)

Raffaele A Calogero

# See Also

oncofuseRun

# **Examples**

#oncofuseInstallation()

oncofuseRun	A function to annotate fusions with Oncofuse. Oncofuse is a naive
	bayesian classifier. Its goal is to identify those fusion sequences with
	higher probability of being driver than passenger events

# Description

A list of fSet objects can be annotated using the Oncofuse java application.

# Usage

```
oncofuseRun(listfSet, tissue=c("EPI","HEM","MES","AVG"), org=c("hg19","hg38"), threads=1, plot=FALSE
```

# Arguments

listfSet	A list of fSets
tissue	Type of tissue in which the fusion was detected. EPI epithelial, HEM ematological, MES mesenchimal, AVG average expression
org	Supported genome assembly version. IMPORTANT: the genome version used for the fusion detection and for the oncofuse analysis need to be same
threads	number of threads that Oncofuse will use

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plot	plotting the expression gain score versus the passenger mutation probability. For

more info please see Shugay et al. Bioinformatics 2013:29,2539

type listfSet a list of fSet objects or coordDf a dataframe of coordinates of fusions in

the format required by Oncofuse

#### Value

The output is a dataframe containing the output of Oncofuse. For more info on Oncofuse please see Shugay et al. Bioinformatics 2013, 29, 2539.

# Author(s)

Raffaele A Calogero

# **Examples**

```
#tmp <- importFusionData("fusionmap", paste(find.package(package="chimera"),"/examples/mcf7.FMFusionReport", se
#installOncofuse()
#of.out <- oncofuseRun(tmp, tissue="EPI")</pre>
```

picardInstallation

A function to download picard-tools

# Description

A function allowing the download of picard-tools in the chimera folder. Picard tools require java

# Usage

picardInstallation()

# Author(s)

Raffaele A Calogero

#### **Examples**

#picardInstallation()

plotCoverage 21

plotCoverage	A function to plot the coverage of a fusion gene	
--------------	--	--

#### **Description**

A function to plot the coverage of a fusion gene.

### Usage

```
\verb|plotCoverage(fset, plot.type=c("exons", "junctions"), junction.spanning=20, fusion.only=FALSE, xlab="linear larger la
```

# **Arguments**

fset	A fSet object
plot.type	exons plot exons coverage as junctions plot coverage of junction between exons
junction.spann	ing
	number of nucleotides located upstream and downstream the junction/fusion location
fusion.only	if TRUE only fusion coverage is plotted
xlab	x-axis label
ylab	y-axis label
main	Plot title
col.box1	color of the box describing the first gene
col.box2	color of the box describing the second gene
ybox.lim	y range defining the height of the box representing the exons

#### Author(s)

Raffaele A Calogero

# See Also

fusionName, chimeraSeqs

```
tmp <- importFusionData("fusionmap", paste(find.package(package="chimera"),"/examples/mcf7.FMFusionReport", sep
fusion.names <- fusionName(tmp)
fusion.names
myset <- tmp[[13]]
trs <- chimeraSeqs(myset, type="transcripts")
myset <- addRNA(myset , trs)
tmp.seq <- fusionRNA(myset)
myset <- addGA(myset, paste(path.package(package="chimera"),"/examples/mcf7_trs_accepted_hits.bam",sep=""))
ga <- fusionGA(myset)
plotCoverage(myset, plot.type="exons", col.box1="red", col.box2="green", ybox.lim=c(-4,-1))</pre>
```

22 removingErrorLine

```
plotCoverage(myset, plot.type="junctions", col.box1="red", col.box2="yellow", ybox.lim=c(-4,-1))
plotCoverage(myset, fusion.only=TRUE, col.box1="red", col.box2="yellow", ybox.lim=c(-4,-1))
```

prettyPrint

A function to represent a list of fSet as a dataframe

#### **Description**

A function reorganizing a list of fSet in a dataframe structure. The dataframe is then saved in a tab delimited file

# Usage

```
prettyPrint(list, filename, fusion.reads=c("all", "spanning"))
```

#### **Arguments**

list a list of fSet object

filename the name of the file in which the dataframe is printed

fusion.reads it allows to extract spanning reads associated to the SeedCount slot or all the

detected reads associate to the RescuedCount

# Author(s)

Raffaele A Calogero

#### **Examples**

```
#tmp <- importFusionData("fusionmap", paste(find.package(package="chimera"),"/examples/mcf7.FMFusionReport", se
#fusion.names <- fusionName(tmp)
#tmp1 <- filterList(tmp, type="fusion.names", fusion.names[c(1,3,7)], parallel=FALSE)
#prettyPrint(tmp1, "tmp1.df.txt", fusion.reads="spanning")</pre>
```

removingErrorLine

A function to remove a line stopping SAM to BAM conversion

#### **Description**

A function to remove a line stopping SAM to BAM conversion

```
removingErrorLine(line.number, filenameIn, filenameOut)
```

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# **Arguments**

line.number line number to be removed

filenameIn input file name filenameOut output file name

#### Value

SAM file without the error line

# Author(s)

Raffaele A Calogero

# **Examples**

```
#removingErrorLine(14680066, "kd2_accepted_hits.sam", "kd2_accepted_hits1.sam")
```

starInstallation

A function to download STAR

# **Description**

A function allowing the download and installation of STAR (Dobin et al. Bioinformatics 2012) in chimera package folder. The function also creates soft links in the user bin folder to allow the call of the above mentioned program.

## Usage

```
starInstallation(binDir, os=c("unix","mac"))
```

# **Arguments**

binDir The user bin folder

os The supported operating systems

# Author(s)

Raffaele A Calogero

```
#starInstallation(binDir="/somewhere/inyourpc/bin", os="mac")
```

24 starRun

starReads

A function to extract reads info from STAR fusion output

# Description

A function producing a GRangeList for the reads information, involved in a fusion event.

# Usage

```
starReads(fusion.report, parallel=FALSE)
```

# **Arguments**

fusion.report STAR fusion output file

parallel option to run a parallel version of the function, default FALSE

#### Author(s)

Raffaele A Calogero

# **Examples**

```
#tmp <- starReads("Chimeric.out.junction", parallel=FALSE)</pre>
```

starRun

A function to generate a bam file for fusions coverage evaluation

#### **Description**

A function mapping reads to a chimera sequence set. The bam produced by this remapping on a putative fusion will be used to plot the coverage data for all the fused constructs. The function assumes that STAR is installed and located in the path.

```
starRun(input1, input2, cores=1, star= "STAR", samtools="samtools", fa, alignment=c("se", "pe"), chimStarRun(input1, input2, cores=1, star= "STAR", samtools="samtools", fa, alignment=c("se", "pe"), chimStarRun(input1, input2, cores=1, star= "STAR", samtools="samtools", fa, alignment=c("se", "pe"), chimStarRun(input1, input2, cores=1, star= "STAR", samtools="samtools", fa, alignment=c("se", "pe"), chimStarRun(input1, input2, cores=1, star= "STAR", samtools="samtools", fa, alignment=c("se", "pe"), chimStarRun(input1, input2, cores=1, star= "starRun(input1, input2, input
```

subreadRun 25

# Arguments

input1	The R1 fastq of a pair-end
input2	The R2 fastq of a pair-end
cores	number of cores to be used by tophat with program name, e.g. /somewhere/tophat
star	full path of STAR with program name, e.g. /somewhere/STAR
samtools	full path of samtools
fa	full path and name of the fasta file of one of the set of fusions of interest, to be used to build the STAR database. The fusion nucleotide sequences was generated with the function chimeraSeqs
alignment	se means that both fastq files from the pair-end run are concatenated, pe means that tophat will use fastq files in pair-end mode

chimSegmentMin STAR fusion parameter, see STAR manual

chimJunctionOverhangMin

STAR fusion parameter, see STAR manual

#### Value

The function create a folder called chimeraDB\_time, where time is the time when the folder was created. STAR output will be located in the folder output\_time, where time is the time when the folder was created. The bam file of interest is accepted\_hits.bam.

#### Author(s)

Raffaele A Calogero

#### See Also

chimeraSeqs

#### **Examples**

 $\#starRun(input1=paste(find.package(package="chimera"),"/examples/mcf7\_sample\_1.fq", sep=""), input2=paste(find.package(package="chimera"),"/examples/mcf7\_sample\_1.fq", sep=""), input2=paste(find.package(package="chimera"),"/examples/mcf7\_sample\_1.fq", sep=""), input2=paste(find.package="chimera"),"/examples/mcf7\_sample\_1.fq", sep=""), input2=paste(find.package="chimera"),"/examples/mcf7\_sample\_1.fq", sep=""), input2=paste(find.package="chimera"),"/examples/mcf7\_sample\_1.fq", sep=""), input2=paste(find.package="chimera"),"/examples/mcf7\_sample\_1.fq", sep=""), input2=paste(find.package="chimera"),"/examples/mcf7\_sample\_1.fq", sep=""), input2=paste(find.package="chimera"),"/examples/mcf7\_sample\_1.fq", sep=""), input2=paste(find.package="chimera"),"/examples/mcf7\_samples/mcf7\_$ 

subreadRun A function to generate a bam file for fusions coverage evaluation

# Description

A function mapping reads to a chimera sequence set. The bam produced by this remapping on a putative fusion will be used to plot the coverage data for all the fused constructs. The function uses Rsubread aligner for MAC and UNIX OS. In case WINDOWS OS Rbowtie is used.

```
subreadRun(ebwt,input1, input2, outfile.prefix="accepted_hits", alignment=c("se","pe"),cores=1)
```

26 supportingReads

### Arguments

ebwt Full path and name of the fasta file of one of the set of fusions of interest, to

be used to build the index database. The fusion nucleotide sequences can be

generated with the function chimeraSeqs

input1 The R1 fastq of a pair-end input2 The R2 fastq of a pair-end

outfile.prefix Prefix of the output bam file. Default is accepted\_hits

alignment se means that both fastq files from the pair-end run are concatenated, pe means

that tophat will use fastq files in pair-end mode

cores Number of cores to be used by the aligner

#### Value

Standard bam file output. The bam file name by default is accepted\_hits.bam.

#### Author(s)

Raffaele A Calogero

#### See Also

chimeraSeqs

#### **Examples**

```
if(require(Rsubread)){
  subreadRun(ebwt=paste(find.package(package="chimera"),"/examples/SULF2_ARFGEF2.fa",sep=""),
    input1=paste(find.package(package="chimera"),"/examples/mcf7_sample_1.fq",sep=""),
    input2=paste(find.package(package="chimera"),"/examples/mcf7_sample_2.fq",sep=""),
    outfile.prefix="accepted_hits", alignment="se", cores=1)
}
```

supportingReads

A function to extract supporting reads values from a list of fSet object

### Description

A function extracting supporting reads values from a list of fSet objects. Please note that not all outputs of supported tools provides both spanning reads, i.e. pair-end reads having one of the two mates spanning over the break point, and encompassing reads, i.e. pair-end reads having the two mates mapping on the exons of the two transcripts involved in the fusion. The presence of both type of reads is mandatory to provide the full number of reads covering the junction region. To know which information are provided by the supported tool please check fSet

```
supportingReads(list, fusion.reads=c("encompassing", "spanning"), parallel=FALSE)
```

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#### **Arguments**

list a list of fSet objects

fusion.reads it allows to extract spanning reads associated to the SeedCount slot or encom-

passing reads associated to the RescuedCount

parallel option to run a parallel version of the function, default FALSE

#### Author(s)

Raffaele A Calogero

#### **Examples**

```
tmp <- importFusionData("fusionmap", paste(find.package(package="chimera"),"/examples/mcf7.FMFusionReport", sep
supporting.reads <- supportingReads(tmp, fusion.reads="spanning")
supporting.reads
```

tophatInstallation

A function to download tophat, bowtie and samtools

# **Description**

A function allowing the download and installation of tophat, bowtie and samtools in chimera package folder. The function also creates soft links in the user bin folder to allow the call of the above mentioned programs.

# Usage

```
tophatInstallation(binDir, os=c("unix","mac"))
```

## **Arguments**

binDir The user bin folder

os The supported operating systems

#### Author(s)

Raffaele A Calogero

```
#tophatInstallation(binDir="/somewhere/inyourpc/bin", os="mac")
```

28 tophatRun

tophatRun	A function to generate a bam file for fusions coverage evaluation

# Description

A function mapping reads to a chimera sequence set. The bam produced by this remapping on a putative fusion will be used to plot the coverage data for all the fused constructs. The function assumes that tophat is installed and located in the path. To run TopHat a softlink to bowtie or bowtie2 need to located in the user bin dir

# Usage

tophatRun(input1, input2, output,cores=1, bowtie= c("bowtie","bowtie2"), tophat= "tophat",ebwt=paste

# **Arguments**

input1	The R1 fastq of a pair-end
input2	The R2 fastq of a pair-end
output	Folder in which tophat will generate the output
cores	number of cores to be used by tophat with program name, e.g. /somewhere/tophat
bowtie	selecting bowtie or bowtie2 aligner
tophat	full path of tophat
ebwt	full path and name of the fasta file of one of the set of fusions of interest, to be used to build the bowtie database. The fusion nucleotide sequences was generated with the function chimeraSeqs
alignment	se means that both fastq files from the pair-end run are concatenated, pe means that tophat will use fastq files in pair-end mode

#### Value

TopHat standard output. The bam file of interest is accepted\_hits.bam. The bam file will be then loaded in the slot fusionsLoc of the fSetSummary object from which fusions were retrieved.

#### Author(s)

Raffaele A Calogero

#### See Also

chimeraSeqs

# **Examples**

#tophatRun(input1=paste(find.package(package="chimera"), "/examples/mcf7\_sample\_1.fq", sep=""), input2=paste(fine)

validateSamFile 29

# Description

A function to validate SAM or BAM files using picard-tools

# Usage

```
validateSamFile(input, output, mode=c("VERBOSE", "SUMMARY"), max.output="100")
```

# Arguments

input SAM/BAM file to be validated

output file name in which to save the validation information

mode Mode of output. Default value: VERBOSE

max.output max number of reported error lines

#### Value

Validation information referring to a SAM/BM file.

# Author(s)

Raffaele A Calogero

# See Also

```
picardInstallation
```

# **Examples**

 $\verb| #validateSamFile(input=paste(find.package(package="chimera"), "/examples/mcf7\_trs\_accepted\_hits.bam", sep=""), or all the package(package="chimera"), "/examples/mcf7\_trs\_accepted\_hits.bam", sep=""), sep=""), sep=""), sep=""), sep=""), sep="">, sep="$ 

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