# Package 'FilterFFPE'

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Description This package finds and filters artificial chimeric reads specifically generated in next-generation sequencing (NGS) process of formalin-fixed paraffin-embedded (FFPE) tissues. These artificial chimeric reads can lead to a large number of false positive structural variation (SV) calls. The required input is an indexed BAM file of a FFPE sample.											
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FilterFFPE-package

FFPE Artificial Chimeric Read Filter for NGS data

#### **Description**

This package finds and filters artificial chimeric reads specifically generated in next-generation sequencing (NGS) process of formalin-fixed paraffin-embedded (FFPE) tissues. These artificial chimeric reads can lead to a large number of false positive structural variation (SV) calls. The required input is an indexed BAM file of a FFPE sample.

#### **Details**

This package was not yet installed at build time.

The next-generation sequencing (NGS) reads from formalin-fixed paraffin-embedded (FFPE) samples contain numerous artifact chimeric reads, which can lead to a large number of false positive structural variation (SV) calls. This package finds and filters these artifact chimeric reads from BAM files of FFPE samples to improve SV calling performance.

Index: This package was not yet installed at build time.

There are three available functions to find and/or filter artifact chimeric reads of FFPE samples:

- 1. findArtifactChimericReads: Find artifact chimeric reads in BAM file of FFPE sample.
- 2. filterBamByReadNames: Filter reads from BAM file by read names.
- 3. FFPEReadFilter: Find and filter artifact chimeric reads in BAM file of FFPE sample.

# Author(s)

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#### See Also

```
FilterFFPE, filterBamByReadNames, FFPEReadFilter
```

#### **Examples**

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Find and filter artifact chimeric reads in BAM file of FFPE sample

#### **Description**

Artifact chimeric reads are enriched in NGS data of FFPE samples, these reads can lead to a large number of false positive SV calls. This function finds and filters these artifact chimeric reads.

#### Usage

```
FFPEReadFilter(file, maxReadsOfSameBreak=2, minMapBase=1, threads=1,
index=file, destination=sub("\\.bam(\\.gz)?", ".FilterFFPE.bam", file),
overwrite=FALSE, FFPEReadsFile=sub("\\.bam(\\.gz)?", ".FFPEReads.txt", file),
dupChimFile=sub("\\.bam(\\.gz)?", ".dupChim.txt", file), filterdupChim=TRUE)
```

# **Arguments**

file Path to the BAM file.

maxReadsOfSameBreak

The maximum allowed number of artifact chimeric reads sharing a false positive breakpoint. If the number of reads sharing the same breakpoint exceeds this number, these reads are not recognized as artifact chimeric reads. Reads marked as PCR or optical duplicates are excluded from the calculation. For paired-end sequencing, a read pair of artifact chimeric fragments may both contain the artifact breakpoints; thereby, the defalut is set to 2.

minMapBase

The minimum required length (bp) of a short complementary mapping for an artifact chimeric read. Artifact chimeric reads are derived from the combination of two single-stranded DNA fragments linked by short reverse complementary regions (SRCR). Reads with SRCR shorter than this length are not recognized as artifact chimeric reads. Note: sequence errors and mutations might influence the detection of the existence and length of SRCR. Suggested range: 0-3. When it is set to 0 or any value below 1, this step will be skipped.

threads Number of threads to use. Multi-threading can speed up the process.

index Path of the index file of the input BAM file.

destination Path of the output filtered BAM file.

overwrite Boolean value indicating whether the destination can be over-written if it already

exists.

FFPEReadsFile Path of the output txt file with artifact chimeric read names.

dupChimFile Path of the output txt file with supplementary reads that are marked as PCR or

optical duplicates.

filterdupChim Filter PCR or optical duplicates of all chimeric reads when set to true. These

reads may contain duplicates of artifact chimeric reads; therefore, it is recom-

mended to also remove these reads.

#### **Details**

The next-generation sequencing (NGS) reads from formalin-fixed paraffin-embedded (FFPE) samples contain numerous artifact chimeric reads, which can lead to a large number of false positive structural variation (SV) calls. This function finds and filters these artifact chimeric reads. An index file is also generated for the created filtered BAM file.

#### Value

The file name of the created destination file.

#### Author(s)

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#### See Also

```
FilterFFPE, findArtifactChimericReads, filterBamByReadNames
```

#### **Examples**

filterBamByReadNames Filter reads from BAM file by read names

# **Description**

Generate filtered BAM file that does not contain reads with the input read names.

#### Usage

```
filterBamByReadNames(file, readsToFilter, index=file,
destination=sub("\\.bam(\\.gz)?", ".FilterFFPE.bam", file), overwrite=FALSE)
```

# **Arguments**

file Path to the input BAM file.

readsToFilter A character vector of read names to filter.

index Path of the index file of the input BAM file.

destination Path of the output filtered BAM file.

overwrite Boolean value indicating whether the destination can be over-written if it already

exists.

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#### **Details**

Generate filtered BAM file that does not contain reads with the input read names, index file is also created

#### Value

The file name of the created destination file.

#### Author(s)

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#### See Also

```
FilterFFPE, findArtifactChimericReads, FFPEReadFilter
```

#### **Examples**

findArtifactChimericReads

Find artifact chimeric reads in BAM file of FFPE sample

# **Description**

Artifact chimeric reads are enriched in NGS data of FFPE samples, these reads can lead to a large number of false positive SV calls. This function finds these artifact chimeric reads.

#### Usage

```
findArtifactChimericReads(file, maxReadsOfSameBreak=2, minMapBase=1,
threads=1, FFPEReadsFile=sub("\\.bam(\\.gz)?", ".FFPEReads.txt", file),
dupChimFile=sub("\\.bam(\\.gz)?", ".dupChim.txt", file))
```

findArtifactChimericReads

#### **Arguments**

file Path to the BAM file. maxReadsOfSameBreak

The maximum allowed number of artifact chimeric reads sharing a false positive breakpoint. If the number of reads sharing the same breakpoint exceeds this number, these reads are not recognized as artifact chimeric reads. Reads marked as PCR or optical duplicates are excluded from the calculation. For paired-end sequencing, a read pair of artifact chimeric fragments may both contain the artifact breakpoints; thereby, the defalut is set to 2.

minMapBase The minimum required length (bp) of a short complementary mapping for an

artifact chimeric read. Artifact chimeric reads are derived from the combination of two single-stranded DNA fragments linked by short reverse complementary regions (SRCR). Reads with SRCR shorter than this length are not recognized as artifact chimeric reads. Note: sequence errors and mutations might influence the detection of the existence and length of SRCR. Suggested range: 0-3. When

it is set to 0 or any value below 1, this step will be skipped.

threads Number of threads to use. Multi-threading can speed up the process.

FFPEReadsFile Path of the output txt file with artifact chimeric read names.

dupChimFile Path of the output txt file with read names of PCR or optical duplicates of all

chimeric reads.

#### **Details**

The next-generation sequencing (NGS) reads from formalin-fixed paraffin-embedded (FFPE) samples contain numerous artifact chimeric reads, which can lead to a large number of false positive structural variation (SV) calls. This function finds the read names of these artifact chimeric reads. To further filter these reads, filterBamByReadNames can be applied.

#### Value

A character vector of artifact chimeric read names.

# Author(s)

Lanying Wei <lanying.wei@uni-muenster.de>

#### See Also

```
FilterFFPE, filterBamByReadNames, FFPEReadFilter
```

#### **Examples**

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