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Abstract

Circulating tumor DNA (ctDNA) containing somatic mutations can be found in blood plasma. This includes DNA fusions, such as the EML4-ALK, which can be an oncogenic driver in non-small cell lung cancer. This is an introduction to the **Genefusiondiscover** package for R, which can be used to evaluate whether EML4-ALK is present in blood plasma.

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1 Introduction

This package was created in order to increase the sensitivity of EML4-ALK detection from commercially available NGS products such as the AVENIO (Roche) pipeline.

Paired-end sequencing of cfDNA generated BAM files can be used as input to discover EML4-ALK variants. This package was developed using position deduplicated BAM files generated with the AVENIO Oncology Analysis Software. These files are made using the AVENIO ctDNA surveillance kit and Illumina Nextseq 500 sequencing. This is a targeted hybridization NGS approach and includes ALK-specific but not EML4-specific probes.

The package includes six functions.

The output of the first function, EML4_ALK_detection(), is used to determine whether EML4-ALK is detected and serves as input for the next four exploratory functions characterizing the EML4-ALK variant. The last function EML4_ALK_analysis() combines the output of the exploratory functions.

To serve as examples, this package includes BAM files representing the EML4-ALK positive cell line H3122 and the EML4-ALK negative cell line, HCC827. # Installation

Use devtools to install Genefusiondiscover.

```
if (!require(devtools)) install.packages('devtools')
library(devtools)

install_github("CTrierMaansson/Genefusiondiscover")
library(Genefusiondiscover)
```

2 Package data

BAM files from the cell lines, H3122 and HCC827, are included in the package and can be used as examples to explore the functions.

```
H3122_bam <- system.file("extdata",

"H3122_EML4.bam",

package = "Genefusiondiscover")

HCC827_bam <- system.file("extdata",

"HCC827_EML4.bam",

package = "Genefusiondiscover")
```

3 Functions

3.1 EML4_ALK_detection()

This function looks for EML4-ALK mate pair reads in the BAM file.

Input:

file

The name of the file which the data are to be read from.

genome

```
character representing the reference genome.
Can either be "hg38" or "hg19".
Default = "hg38".
```

mates

```
integer, the minimum number EML4-ALK mate pairs needed to be detected in order to call a variant. Default = 2.
```

Output:

If EML4-ALK is detected, a data.frame with soft-clipped reads representing EML4-ALK is returned. Otherwise "No EML4-ALK was detected" is returned.

Examples:

```
head(EML4_ALK_detection(file = H3122_bam,
       genome = "hg38",
       mates = 2))
mate position cigar
#> 726 29223691 42299657 94M2S
#> 727 29223375 42299657 94M2S
#> 728 29223479 42299657 94M2S
#> 729 29223686 42299657 94M2S
#> 731 29223636 42299657 94M2S
#> 733 29223687 42299657 94M2S
```

3.2 EML4_sequence()

This function identifies the basepairs leading up to the EML4 breakpoint.

Input:

reads

```
data.frame returned by EML4_ALK_detection().
```

basepairs

```
integer, number of basepairs identified from the EML4-ALK fusion. Default = 20.
```

Output:

If EML4-ALK is detected, a table of identified EML4 basepairs is returned, with the number of corresponding reads for each sequence. Otherwise "No EML4-ALK was detected" is returned.

Examples:

3.3 ALK_sequence()

This function identifies the basepairs following the ALK breakpoint.

Input:

reads

```
data.frame returned by EML4_ALK_detection().
```

basepairs

```
integer, number of basepairs identified from the EML4-ALK fusion.

Default = 20.
```

Output:

If EML4-ALK is detected, a table of identified ALK basepairs is returned, with the number of corresponding reads for each sequence. Otherwise "No EML4-ALK was detected" is returned.

Examples:

```
ALK_sequence(EML4_ALK_detection(file = H3122_bam,
                                genome = hg38,
                                mates = 2),
                basepairs = 20)
#> ALK_seg
#> CAGAATTTTAGCTTTGCAAT
                            CGGATTTTTAGCTTT CGGATTTTTAGCTTTTCATT
                                          1
                                                               2
                                          CT
#> CGGAATTTTAGCTTTGCATT
                                                              CTG
#>
                     1
                                          8
                                                                3
                   CTGA
                                       CTGAT
#>
                                                        CTGATTTT
                   11
                                       16
                                                               5
#>
             CTGATTTTT
                                 CTGATTTTTA
                                                     CTGATTTTTAG
#> CTGATTTTTAGATTTGCATT
                                CTGATTTTTAGC
                                                    CTGATTTTTAGCT
                                         14
#>
                             CTGATTTTTAGCTTT
        CTGATTTTTAGCTT
                                                 CTGATTTTTAGCTTTG
#>
                    10
                                          3
#>
     CTGATTTTTAGCTTTGC
                         CTGATTTTTAGCTTTGCA CTGATTTTTAGCTTTGCAT
#>
                     7
                                          8
#> CTGATTTTTAGCTTTGCATT CTGATTTTTAGCTTTGCAAT
                                                 CTGATTTTTAGCTTTT
                    71
                                          1
#> CTGATTTTTAGCTTTTCATA
                                 CTGATTTTTAT
                                                 CTGATTTTTATCTTTG
#>
                     1
                                          1
#> CTGATTTTTATCTTTGCATT CTGATTTTTATCTTTTGATT CTGTGTTTTTAGATTTGCATT
                      2
                                          1
                                                                1
#> CTGTTTTTTATCTTTGCAAT
                                       CTGAA CTTATTTTTATCTTTGCATT
```

3.4 break_position()

This function identifies the genomic position in EML4 where the breakpoint has happened.

Input:

reads

```
data.frame returned by EML4_ALK_detection().
```

Output:

If EML4-ALK is detected, a table of genomic positions is returned with the number of corresponding reads for each sequence. Otherwise "No EML4-ALK was detected" is returned.

Examples:

3.5 break_position_depth()

This function identifies the read depth at the basepair before the breakpoint in EML4.

Input:

file

The name of the file which the data are to be read from.

reads

```
data.frame returned by EML4_ALK_detection().
```

Output:

If EML4-ALK is detected a single integer corresponding to the read depth at the breakpoint is returned. Otherwise "No EML4-ALK was detected" is returned

Examples:

3.6 EML4_ALK_analysis()

This functions collects the results from the other functions of the package.

Input:

file

The name of the file which the data are to be read from.

genome

```
character representing the reference genome.
Can be either "hg38" or "hg19".
Default = "hg38".
```

mates

```
integer, the minimum number EML4-ALK mate pairs needed to be detected in order to call a variant. Default = 2.
```

basepairs

```
integer, number of basepairs identified from the EML4-ALK fusion.
Default = 20.
```

Output:

A list object with clipped_reads corresponding to EML4_ALK_detection(), last_EML4 corresponding to EML4_sequence(), first_ALK corresponding to ALK_sequence(), breakpoint corresponding to break_position(), and read_depth corresponding to break_position_depth().

Examples:

```
H3122_results <- EML4_ALK_analysis(file = H3122_bam,
                       genome = hg38,
                       mates = 2,
                       basepairs = 20)
HCC827_results <- EML4_ALK_analysis(file = HCC827_bam,</pre>
                       genome = "hg38",
                       mates = 2,
                       basepairs = 20)
head(H3122_results$clipped_reads)
mate position cigar
#> 726 29223691 42299657 94M2S
#> 727 29223375 42299657 94M2S
#> 728 29223479 42299657 94M2S
#> 729 29223686 42299657 94M2S
#> 731 29223636 42299657 94M2S
#> 733 29223687 42299657 94M2S
H3122_results$last_EML4
#> EML4_seq
#> CCAGGCTGGAGTGCAGTGGT GGAGTGCAGTGGTGTTT TCAGGCTGGAGTGCAGTGGT
                           1
             201
H3122_results$first_ALK
#> ALK_seg
#> CAGAATTTTAGCTTTGCAAT
                  CGGATTTTTAGCTTT CGGATTTTTAGCTTTTCATT
```

```
TTGCATT CT CTG

1 8 3

CTGA CTGAT CTGATTTT

11 16 5

GATTTTT CTGATTTTTA CTGATTTTTAG
#> CGGAATTTTAGCTTTGCATT
#> 1
#> CTGA
          11
        CTGATTTTT
                         3
#> CTGATTTTTAGATTTGCATT CTGATTTTTAGC CTGATTTTTAGCT #> 1 14 10
#>
    CTGATTTTTAGCTT CTGATTTTTAGCTTT CTGATTTTTAGCTTTG
      10
#> CTGATTTTTAGCTTTGC CTGATTTTTAGCTTTGCA CTGATTTTTAGCTTTGCAT
#> 7 8 1
#> CTGATTTTTAGCTTTGCATT CTGATTTTTAGCTTTTGCAAT CTGATTTTTAGCTTTT
#> 71 1
#> CTGATTTTTAGCTTTTCATA CTGATTTTTAT CTGATTTTTATCTTTG
#> 1 1 2
                     1 2
#> CTGATTTTTATCTTTGCATT CTGATTTTTATCTTTTGATT CTGTGTTTTAGATTTGCATT
#> 2
#> 2 1 1
#> CTGTTTTTATCTTTGCAAT CTGAA CTTATTTTATCTTTGCATT
#> 1
                           1 1
#> TTAGCTTTG
H3122_results$breakpoint
#> break_pos
#> 42299750 42299757
#> 202 1
H3122_results$read_depth
#> [1] 251
HCC827_results
#> [1] "No EML4-ALK was detected"
```

4 Session info

```
#> - Session info -----
#> setting value
#> version R version 4.2.1 (2022-06-23 ucrt)
#> os Windows 10 x64 (build 22000)
\# system x86_64, mingw32
#> ui RTerm
#> language (EN)
#> collate Danish_Denmark.utf8
#> ctype Danish_Denmark.utf8
#> tz Europe/Paris
#> date 2022-07-27
#> pandoc 2.17.1.1 @ C:/Program Files/RStudio/bin/quarto/bin/ (via rmarkdown)
#> - Packages ------
* 2.4.4 2022-07-20 [1] CRAN (R 4.2.1)
#> Genefusiondiscover * 0.99.0 2022-07-27 [1] Github (CTrierMaansson/Genefusiondiscover@8e86567)
#> [1] C:/Users/Christoffer/AppData/Local/R/win-library/4.2
#> [2] C:/Program Files/R/R-4.2.1/library
#> -----
```