Package 'microhaplot'

October 13, 2022

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Type Package
Title Microhaplotype Constructor and Visualizer
Version 1.0.1
Description A downstream bioinformatics tool to construct and assist curation of microhaplotypes from short read sequences.
Depends R (>= $3.1.2$)
Encoding UTF-8
License GPL-3
LazyData TRUE
Imports DT (>= 0.1), dplyr (>= 0.4.3), ggplot2 (>= 2.1.0), grid (>= 3.1.2), gtools (>= 3.5.0), magrittr (>= 1.5), scales (>= 0.4.0), shiny (>= 0.13.2), shinyBS (>= 0.61), tidyr (>= 0.4.1), shinyWidgets (>= 0.4.3), ggiraph (>= 0.6.0)
<pre>URL https://github.com/ngthomas/microhaplot</pre>
BugReports https://github.com/ngthomas/microhaplot/issues
RoxygenNote 6.1.1
Suggests knitr, rmarkdown
VignetteBuilder knitr
NeedsCompilation no
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Repository CRAN
Date/Publication 2019-10-03 07:40:05 UTC
R topics documented:
mvShinyHaplot
Index

2 prepHaplotFiles

mvShinyHaplot

Transfer a copy of microhaplot app.

Description

Moves shiny microhaplot app to a different directory

Usage

```
mvShinyHaplot(path)
```

Arguments

path

string. directory path. Required

Value

a logical value of whether file.copy is successfully transferred Shiny app to its new directory

Examples

```
mvShinyHaplot(tempdir())
```

prepHaplotFiles

Extracts haplotype from alignment reads.

Description

The function microhaplot extracts haplotype from sequence alignment files through perl script hapture and returns a summary table of the read depth and read quality associate with haplotype.

Usage

```
prepHaplotFiles(run.label, sam.path, label.path, vcf.path,
  out.path = tempdir(), add.filter = FALSE, app.path = tempdir(),
  n.jobs = 1)
```

prepHaplotFiles 3

Arguments

character vector. Run label to be used to display in haPLOType. Required
string. Directory path folder containing all sequence alignment files (SAM). Required
string. Label file path. This customized label file is a tab-separate file that contains entries of SAM file name, individual ID, and group label. Required
string. VCF file path. Required
string. Optional. If not specified, the intermediate files are created under TEMPDIR, with the assumption that directory is granted for written permission.
boolean. Optional. If true, this removes any haplotype with unknown and deletion alignment characters i.e. "*" and "_", removes any locus with large number of haplotypes ($\# > 40$) , and remove any locus with fewer than half of the total individuals.
string. Path to shiny haPLOType app. Optional. If not specified, the path is default to TEMPDIR.
positive integer. Number of SAM files to be parallel processed. Optional. This multithread is only available for non Window OS. Recommend two times the number of processors/core.

Value

This function returns a dataframe of 9 columns i.e group, id, locus, haplotype, depth, sum of Phred score, max of Phred score, allele balance and haplotype rank from highest to lowest read depth. This dataframe will also be saved in out.path.

Examples

4 runShinyHaplot

```
sam.path = sam.path,
out.path = tempdir(),
label.path = label.path,
vcf.path = vcf.path,
app.path = app.path)
}else {
message("Perl version is outdated. Must >= 5.014.")}
```

runShinyHaplot

Run shiny microhaplot

Description

Run shiny microhaplot app

Usage

```
runShinyHaplot(path = system.file("shiny", "microhaplot", package =
   "microhaplot"))
```

Arguments

path

Path to shiny microhaplot app. Optional. If not specified, the path is default to local app path.

Value

Runs shiny microhaplot application via shiny::runApp which typically doesn't return; interrupt R to stop the application (usually by pressing Ctrl+C or Esc).

Examples

```
if(interactive()){
runShinyHaplot()
}
```

Index

```
mvShinyHaplot, 2
prepHaplotFiles, 2
runShinyHaplot, 4
```