Package: neutralitytestr (via r-universe)

November 1, 2024

Title Test for a Neutral Evolutionary Model in Cancer Sequencing Data

Version 0.0.3

Description Package takes frequencies of mutations as reported by high throughput sequencing data from cancer and fits a theoretical neutral model of tumour evolution. Package outputs summary statistics and contains code for plotting the data and model fits. See Williams et al 2016 <doi:10.1038 ng.3489=""> and Williams et al 2017 <doi:10.1101 096305=""> for further details of the method.</doi:10.1101></doi:10.1038>								
Depends R (>= 3.4)								
License MIT + file LICENSE								
Encoding UTF-8								
LazyData true								
Imports dplyr, ggplot2, scales, pracma, ggpmisc, cowplot								
Suggests knitr, rmarkdown, testthat								
VignetteBuilder knitr								
<pre>URL https://github.com/marcjwilliams1/neutralitytestr</pre>								
BugReports https://github.com/marcjwilliams1/neutralitytestr/issues								
RoxygenNote 7.1.1								
Repository https://marcjwilliams1.r-universe.dev								
RemoteUrl https://github.com/marcjwilliams1/neutralitytestr								
RemoteRef HEAD								
RemoteSha 86911335a5b22c161adcd6184665faaf0386a95d								
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 lsq_plot

Plot cumulative distribution lsq_plot Plots the cumulative distribution of the data as well as the best fit linear model line.

Description

Plot cumulative distribution lsq_plot Plots the cumulative distribution of the data as well as the best fit linear model line.

Usage

```
lsq_plot(object)
```

Arguments

object

neutrality test object

Value

ggplot object.

Examples

```
lsq_plot(neutralitytest(VAFselection, fmin = 0.1, fmax = 0.25))
```

neutralitytest

Testing for neutrality on cancer sequencing data

Description

neutralitytest returns a neutralitytest object which contains the result of various test statistics to test for neutrality as described in Williams et al. Nature Genetics 2018. WARNING: This package has been superseded by MOBSTER, see Caravagna et al. Nature Genetics 2020.

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Usage

```
neutralitytest(
  VAF,
  fmin = 0.1,
  fmax = 0.25,
  read_depth = NULL,
  rho = 0,
  cellularity = 1,
  ploidy = 2
)
```

Arguments

VAF Vector of variant allele frequencies (VAFs) from a deep sequencing experiment,

numbers should be between 0 and 1

fmin Minimum VAF of integration range, default is 0.1

fmax Maximum VAF of integration range, default is 0.25

read_depth Read depth of sample, if this is specified it will be used to calculate an approp-

tiate integration range. default is NULL in which case the default or inputted

fmin and fmax will be used.

rho Overdispersion of sample if known, default is 0.0. Will be used to calculate

integration range if read_depth != NULL

cellularity Cellularity of sample, default is 1.0. Will be used to calculate integration range

if read_depth != NULL

ploidy Ploidy of the genome, default is 2. Ideally mutations should be filtered for this

ploidy before running the test. Will be used to calculate integration range if

read_depth != NULL

Value

neutralitytest object which contains test statistics which tests if the sequencing data is consistent a neutral evolutionary model. Test statistics are area between theoretical and empirical curves, kolmogorov distance, mean distance and R^2 statistics from linear model fit. Also returns an estimate of the mutation rate per tumour tumour doubling, the raw VAFs and cumulative distribution

Examples

```
neutralitytest(runif(100))
neutralitytest(VAFselection, fmin = 0.1, fmax = 0.25)
neutralitytest(VAFneutral, read_depth = 100.0, cellularity = 0.8)
```

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neutralitytestr

neutralitytestr package

Description

Package to test a neutral evolutionary model on deep sequencing data.

Details

See the README on GitHub

normalized_plot

Plot normalized cumulative distribution normalized_plot Plots the (normalized) cumulative distribution of the data as well as the theoretical expectation from a neutral evolutionary model.

Description

Plot normalized cumulative distribution normalized_plot Plots the (normalized) cumulative distribution of the data as well as the theoretical expectation from a neutral evolutionary model.

Usage

```
normalized_plot(object)
```

Arguments

object

neutrality test object

Value

ggplot object.

Examples

```
normalized_plot(neutralitytest(VAFselection, fmin = 0.1, fmax = 0.25))
```

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plot_all	Plot all plots in the package and make composite figure. plot_all Plots histogram, linear model best fit plot and normalized plot and plot and makes composite figure.

Description

Plot all plots in the package and make composite figure. plot_all Plots histogram, linear model best fit plot and normalized plot and plot and makes composite figure.

Usage

```
plot_all(object)
```

Arguments

object

neutrality test object

Value

ggplot object.

Examples

```
plot_all(neutralitytest(VAFselection, fmin = 0.1, fmax = 0.25))
```

VAFneutral

Synthetic sequencing data generated from a evolutionary based cancer simulation.

Description

This data is generated from a neutral evolutionary model where all subclonal mutations are neutral.

Usage

VAFneutral

Format

A vector with variant allele frequencies (VAFs) ranging from 0 to 1

Source

Generated using cancer sequencing simulation https://github.com/marcjwilliams1/CancerSeqSim.jl

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VAFselection	Synthetic sequencing data generated from a evolutionary based cancer simulation.

Description

This data is generated from an evolutionary model where there is on subclonal population and all other mutations are neutral passengers.

Usage

VAFselection

Format

A vector with variant allele frequencies (VAFs) ranging from 0 to 1

Source

Generated using cancer sequencing simulation https://github.com/marcjwilliams1/CancerSeqSim.jl

vaf_histogram	Plot VAF histogram vaf_histogram Plots a histogram of the variant
	allele frequencies.

Description

Plot VAF histogram vaf_histogram Plots a histogram of the variant allele frequencies.

Usage

```
vaf_histogram(object)
```

Arguments

object

neutrality test object

Value

ggplot object.

Examples

```
vaf_histogram(neutralitytest(VAFselection, fmin = 0.1, fmax = 0.25))
```

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