

Package ‘junctions’

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Type Package

Title The Breakdown of Genomic Ancestry Blocks in Hybrid Lineages

Version 2.1.3

Description Individual based simulations of hybridizing populations, where the accumulation of junctions is tracked. Furthermore, mathematical equations are provided to verify simulation outcomes. Both simulations and mathematical equations are based on Janzen (2018, <[doi:10.1101/058107](https://doi.org/10.1101/058107)>) and Janzen (2022, <[doi:10.1111/1755-0998.13519](https://doi.org/10.1111/1755-0998.13519)>).

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URL <https://thijsjanzen.github.io/junctions/>

BugReports <https://github.com/thijsjanzen/junctions/issues>

Depends RcppParallel (>= 5.0.0)

Imports nloptr, Rcpp, tibble

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LinkingTo nloptr, Rcpp, RcppParallel

VignetteBuilder knitr

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junctions-package	<i>Extending The Theory of Junctions</i>
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Description

The theory of junctions is extended by this package by including the effect of a finite number of recombination sites along the chromosome. The package provides functions to calculate the estimated number of junctions, depending on the time since the onset of hybridization, population size, number of recombination sites, initial heterozygosity and the number of crossovers per meiosis.

Details

This package provides individual based simulations in order to simulate the accumulation of junctions over time, both for chromosomes with a finite and an infinite number of recombination sites. Furthermore, the package provides mathematical tools to verify the outcomes of the individual based simulations.

Update version 2.1.1 : fixed typo in one of the examples

Update version 2.1.0 : updated `tbb::task_scheduler_init` to `tbb::global_control`

Update version 2.0.2 : simplified some tests

Update version 2.0 : merged many functions with similar functionality, added vignette that provides overview of all functionality.

Update version 1.9 : added c++ versions of the unphased and phased likelihoods.

Update version 1.8 : added multithreading using the TBB library.

Update version 1.7 : further improved the recombination function following Hanno Hildenbrandt's suggestions

Update version 1.6 : improved the recombination function to run twice as fast

Update version 1.5.1: added option to track the true number of junctions

Update version 1.5: added support for inferring the time since admixture based on phased and unphased data. Also included are simulation functions to simulate appropriate data (e.g. phased and unphased).

Update version 1.4: added support for estimating the number of junctions, and simulating the number of junctions, under a backcrossing scheme, using the code supplied in Lavretsky et al. 2019.

Update version 1.3: added support for estimating the time since admixture using unphased data.

Update version 1.3: added individual based simulations returning phased and unphased data.

Update version 1.3: Updated entire package to Roxygen.

Update version 1.2: added support for estimating the expected number of junctions for arbitrarily distributed markers.

Update version 1.1: updated underlying random number generator for picking recombination sites. The previous generator had limited precision, which could generate duplicate recombination sites. This update fixes that

Author(s)

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References

Janzen, T. , Nolte, A. W. and Traulsen, A. (2018), The breakdown of genomic ancestry blocks in hybrid lineages given a finite number of recombination sites. *Evolution*, 72: 735-750. doi:10.1111/evo.13436

Lavretsky, P, Janzen, T. and McCracken, KG. (2019) Identifying hybrids & the genomics of hybridization: Mallards & American black ducks of Eastern North America. *Ecology and Evolution* 9: 3470-3490. doi:10.1002/ece3.4981

calculate_mat	<i>Function to calculate the maximum accurate time</i>
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Description

Function that calculates the maximum time after hybridization after which the number of junctions can still be reliably used to estimate the onset of hybridization. This is following equation 15 in Janzen et al. 2018.

Usage

```
calculate_mat(N = Inf, R = Inf, H_0 = 0.5, C = 1)
```

Arguments

N	Population Size
R	Number of genetic markers
H_0	Frequency of heterozygosity at $t = 0$
C	Mean number of crossovers per meiosis (e.g. size in Morgan of the chromosome)

Value

The maximum accurate time

Examples

```
calculate_mat(N = Inf, R = 1000, H_0 = 0.5, C = 1)
```

calc_k	<i>Calculate the limit of the number of junctions</i>
--------	---

Description

Calculate the average number of junctions after an infinite number of generations, provided information on the initial heterozygosity, population size and the number of generations.

Usage

```
calc_k(N = Inf, R = Inf, H_0 = 0.5, C = 1)
```

Arguments

N	population size
R	number of markers
H_0	initial heterozygosity (at the time of admixture)
C	Mean number of crossovers per meiosis (e.g. size in Morgan of the chromosome)

Value

The number of junctions for at time = infinity

Examples

```
k <- calc_k(N = 100, R = 1000, H_0 = 0.5, C = 1)
```

estimate_time	<i>Estimate the time since the onset of hybridization, using the number of junctions</i>
---------------	--

Description

Estimate the time since the onset of hybridization, following equation 14 in Janzen et al. 2018

Usage

```
estimate_time(J = NA, N = Inf, R = Inf, H_0 = 0.5, C = 1)
```

Arguments

J	The observed number of junctions
N	Population Size
R	Number of genetic markers
H_0	Frequency of heterozygosity at t = 0
C	Mean number of crossovers per meiosis (e.g. size in Morgan of the chromosome)

Value

The number of generations passed since the onset of hybridization

Examples

```
J <- number_of_junctions(N = 100, R = 1000, H_0 = 0.5, C = 1, t = 200)
estimate_time(J = J, N = 100, R = 1000, H_0 = 0.5, C = 1)
# should be 200 again
```

`estimate_time_diploid` *estimates the time since admixture, given diploid ancestry data.*

Description

Calculates the time since admixture, given unphased ancestry data.

Usage

```
estimate_time_diploid(
  ancestry_information,
  analysis_type = "individuals",
  phased = FALSE,
  pop_size = 1000,
  freq_ancestor_1 = 0.5,
  lower_lim = 2,
  upper_lim = 2000,
  num_threads = 1,
  verbose = FALSE
)
```

Arguments

<code>ancestry_information</code>	a matrix with five columns: column 1) indicator of individual, column 2) indicator of chromosome, 3) location of marker in Morgan, 4) ancestry at chromosome 5) ancestry at chromosome 2.
<code>analysis_type</code>	how should the data be broken down? there are multiple options: "individuals" - time is inferred for each individual separately, grouping all chromosomes together that belong to the same individual. "chromosomes" - time is inferred for each chromosome separately, grouping chromosomes together belonging from separate individuals. "separate" - time is inferred for each chromosome from each individual separately, "all" - time is inferred jointly for all chromosomes and individuals, grouping all chromosomes and individuals together.
<code>phased</code>	is the data phased?
<code>pop_size</code>	population size
<code>freq_ancestor_1</code>	Frequency of ancestor 1 at $t = 0$
<code>lower_lim</code>	lower limit of the optimization algorithm. Increase if the expected admixture time is relatively ancient
<code>upper_lim</code>	upper limit of the optimization algorithm. If set too large, recent admixture events can be overlooked - best to set as low as possible.
<code>num_threads</code>	<code>num_threads</code> , default is all threads. 5 threads is recommended.
<code>verbose</code>	display intermediate output? Default = FALSE

estimate_time_haploid *estimate time using likelihood for a single chromosome*

Description

Estimate the time since the onset of hybridization, for a haploid genome

Usage

```
estimate_time_haploid(
  ancestry_matrix,
  N = 1000,
  freq_ancestor_1 = 0.5,
  lower_lim = 2,
  upper_lim = 1000,
  verbose = FALSE
)
```

Arguments

ancestry_matrix	matrix with 3 columns, column 1 = chromosome, column 2 = location in Morgan, column 3 = ancestry.
N	Population Size
freq_ancestor_1	Frequency of ancestor 1 at $t = 0$
lower_lim	lower limit of the optimization algorithm. Increase if the expected admixture time is relatively ancient
upper_lim	upper limit of the optimization algorithm. If set too large, recent admixture events can be overlooked - best to set as low as possible.
verbose	return verbose output

Value

The number of generations passed since the onset of hybridization

estimate_time_one_chrom

Estimate the time since the onset of hybridization, using the observed number of junctions, taking into account the distribution of markers on a single chromosome

Description

Estimate the time since the onset of hybridization, following equation 1 in Janzen et al. unpublished

Usage

```
estimate_time_one_chrom(
  J = NA,
  N = Inf,
  H_0 = 0.5,
  marker_distribution = NA,
  lower_lim = 2,
  upper_lim = 1000
)
```

Arguments

J	The observed number of junctions
N	Population Size
H_0	Frequency of heterozygosity at $t = 0$
marker_distribution	A vector containing the position of all markers in Morgan.
lower_lim	lower limit of the optimization algorithm. Increase if the expected admixture time is relatively ancient
upper_lim	upper limit of the optimization algorithm. If set too large, recent admixture events can be overlooked - best to set as low as possible.

Value

The number of generations passed since the onset of hybridization

Examples

```
markers <- seq(from = 0, to = 1, length.out = 100)
J <- number_of_junctions_markers(N = 100, H_0 = 0.5, t = 200,
  marker_distribution = markers)
estimate_time_one_chrom(J = J,
  N = 100,
  H_0 = 0.5,
  marker_distribution = markers) #should be 200 again
```

log_likelihood_diploid

calculate the log likelihood of observing diploid ancestry data.

Description

Calculates the log likelihood of observing the phased data, given the population size, initial heterozygosity and time since admixture

Usage

```
log_likelihood_diploid(
  local_anc_matrix,
  pop_size,
  freq_ancestor_1 = 0.5,
  t,
  phased = FALSE,
  num_threads = 1
)
```

Arguments

local_anc_matrix	a matrix with four columns: column 1) chromosome indicator, 2) location of marker in Morgan on respective chromosome 3) ancestry at chromosome 4) ancestry at chromosome 2.
pop_size	population size
freq_ancestor_1	Frequency of ancestor 1 at t = 0
t	time since admixture
phased	is the data phased or not? default is false.
num_threads	number of threads, default is one thread. Set to -1 to use all available threads.

Value

log likelihood

log_likelihood_haploid

log likelihood of the time since admixture for a haploid genome

Description

log likelihood of the time since admixture for a set of single chromosomes (for ex. in Yeast).

Usage

```
log_likelihood_haploid(ancestry_matrix, N = 1000, freq_ancestor_1 = 0.5, t = 2)
```

Arguments

ancestry_matrix	matrix with 3 columns, column 1 = chromosome, column 2 = location in Morgan, column 3 = ancestry.
N	Population Size
freq_ancestor_1	Frequency of ancestor 1 at t = 0
t	time since admixture

Value

loglikelihood

number_of_junctions	<i>Calculate the average number of junctions</i>
---------------------	--

Description

Calculate the average number of junctions in a single chromosome after t generations, provided information on the initial heterozygosity, population size and the number of generations.

Usage

```
number_of_junctions(N = Inf, R = Inf, H_0 = 0.5, C = 1, t = 100)
```

Arguments

N	Population Size
R	Number of genetic markers
H_0	Frequency of heterozygosity at t = 0
C	Mean number of crossovers per meiosis (e.g. size in Morgan of the chromosome)
t	Time since admixture

Value

Estimated number of junctions at time t

Examples

```
jt <- number_of_junctions(N = 100, R = 1000, H_0 = 0.5, C = 1, t = 1000)
jt2 <- number_of_junctions(N = 100, R = 1000, H_0 = 0.5, C = 1, t = 0:1000)
```

number_of_junctions_backcross

Calculate the average number of junctions during backcrossing

Description

Calculate the expected number of junctions after t generations, in a backcrossing mating scheme.

Usage

```
number_of_junctions_backcross(H_0 = 0.5, C = 1, t = 100)
```

Arguments

H_0	Frequency of heterozygosity at $t = 0$
C	Mean number of crossovers per meiosis (e.g. size in Morgan of the chromosome)
t	Time since admixture

Value

Estimated number of junctions at time t

Examples

```
jt <- number_of_junctions_backcross(H_0 = 0.1, C = 1, t = 5)
```

number_of_junctions_di

Calculate the expected number of junctions between two markers separated by a given amount of recombination

Description

Calculate the expected number of junctions after t generations, provided information on the initial heterozygosity, population size, the number of generations since the onset of admixture and the distance between two markers.

Usage

```
number_of_junctions_di(N = Inf, H_0 = 0.5, t = 100, di = 1e-06)
```

Arguments

N	Population Size
H_0	Frequency of heterozygosity at $t = 0$
t	Time since admixture
di	Distance between two markers in Morgan

Value

Estimated number of junctions at time t

Examples

```
number_of_junctions_di(N = 100, H_0 = 0.5, t = 1000, di = 0.01)
```

```
number_of_junctions_markers
```

*Calculate the expected total number of junctions in a chromosome,
given the distribution of markers*

Description

Calculate the expected number of junctions after t generations, provided information on the initial heterozygosity, population size, the number of generations since the onset of admixture and the distribution of markers.

Usage

```
number_of_junctions_markers(  
  N = Inf,  
  H_0 = 0.5,  
  t = 100,  
  marker_distribution = NA  
)
```

Arguments

N	Population Size
H_0	Frequency of heterozygosity at $t = 0$
t	Time since admixture
marker_distribution	A vector containing the position of all markers in Morgan.

Value

Estimated number of observed junctions at time t

Examples

```

markers <- seq(from = 0, to = 1, length.out = 1000)
jt <- number_of_junctions_markers(N = 100,
                                H_0 = 0.5,
                                t = 1000,
                                marker_distribution = markers)
random_markers <- sort(runif(1000, 0, 1))
jt2 <- number_of_junctions_markers(N = 100,
                                H_0 = 0.5,
                                t = 1000,
                                marker_distribution = random_markers)

```

sim_backcrossing	<i>Function to simulate data using a back crossing scheme</i>
------------------	---

Description

Individual based simulation of the accumulation of junctions, under a back crossing scheme

Usage

```

sim_backcrossing(
  population_size = 100,
  freq_ancestor_1 = 0.5,
  total_runtime = 5,
  size_in_morgan = 1,
  number_of_markers = 100,
  seed = 6,
  time_points = -1
)

```

Arguments

population_size	Population size
freq_ancestor_1	Frequency of ancestor 1 at t = 0
total_runtime	Number of generations to simulate
size_in_morgan	Mean number of crossovers per meiosis (e.g. size in Morgan of the chromosome)
number_of_markers	number of molecular markers
seed	Seed of the pseudo-random number generator
time_points	vector with time points at which local ancestry has to be recorded to be returned at the end of the simulation. If left at -1, ancestry is recorded at every generation (computationally heavy).

Value

List with five entries: average_junctions: average number of junctions over time, detected_junctions: average number of detected junctions, given the markers. markers: vector with the locations of the molecular markers, junction_distribution: distribution of junctions per time step average_heterozygosity: average heterozygosity.

Examples

```
sim_backcrossing(population_size = 100,
                  total_runtime = 5,
                  size_in_morgan = 1,
                  number_of_markers = 100,
                  seed = 6,
                  time_points = 1:5)
```

sim_fin_chrom	<i>Individual Based Simulation of the accumulation of junctions</i>
---------------	---

Description

Individual based simulation of the accumulation of junctions for a chromosome with regularly distributed markers.

Usage

```
sim_fin_chrom(
  pop_size = 100,
  freq_ancestor_1 = 0.5,
  total_runtime = 100,
  morgan = 1,
  seed = 42,
  R = 100
)
```

Arguments

pop_size	Population Size
freq_ancestor_1	Frequency of ancestor 1 at t = 0
total_runtime	Maximum time after which the simulation is to be stopped
morgan	Mean number of crossovers per meiosis (e.g. size in Morgan of the chromosome)
seed	Seed of the pseudo-random number generator
R	Number of regularly distributed markers

Value

avgJunctions vector of the average number of junctions at time = [0, total_runtime]

Examples

```
sim_inf_chrom(pop_size = 100, freq_ancestor_1 = 0.5,
              total_runtime = 10, morgan = 1, seed = 42,
              R = 100)
```

sim_inf_chrom	<i>Individual Based Simulation of the accumulation of junctions</i>
---------------	---

Description

Individual based simulation of the accumulation of junctions for a chromosome with an infinite number of recombination sites.

Usage

```
sim_inf_chrom(
  pop_size = 100,
  freq_ancestor_1 = 0.5,
  total_runtime = 100,
  morgan = 1,
  markers = -1,
  seed = 42
)
```

Arguments

pop_size	Population Size
freq_ancestor_1	Frequency of ancestor 1 at t = 0
total_runtime	Maximum time after which the simulation is to be stopped
morgan	Mean number of crossovers per meiosis (e.g. size in Morgan of the chromosome)
markers	The number of genetic markers superimposed on the chromosome. If markers is set to -1, no markers are superimposed (faster simulation)
seed	Seed of the pseudo-random number generator

Value

avgJunctions vector of the average number of junctions at time = [0, total_runtime]

Examples

```
v <- sim_inf_chrom(pop_size = 100, freq_ancestor_1 = 0.5,
                  total_runtime = 10, morgan = 1, markers = 100,
                  seed = 42)
plot(v$avgJunctions, type = "l", xlab = "Generations",
     ylab = "Number of Junctions", main = "Example Infinite Chromosome")
lines(v$detectedJunctions, col = "blue")
legend("bottomright", c("Real number", "Number detected"),
      lty = 1, col = c("black", "blue"))
```

sim_phased_unphased	<i>Individual Based Simulation of the accumulation of junctions</i>
---------------------	---

Description

Individual based simulation of the accumulation of junctions, returning phased and unphased data. Ancestry on both chromosomes of 10 randomly sampled individuals per generations is returned.

Usage

```
sim_phased_unphased(
  pop_size = 100,
  freq_ancestor_1 = 0.5,
  total_runtime = 100,
  size_in_morgan = 1,
  markers = 100,
  time_points = -1,
  num_threads = 1,
  verbose = FALSE,
  record_true_junctions = FALSE,
  num_indiv_sampled = 10,
  coverage = 1,
  error_rate = 0
)
```

Arguments

pop_size	Population Size
freq_ancestor_1	Frequency of ancestor 1 at t = 0
total_runtime	Maximum time after which the simulation is to be stopped
size_in_morgan	Mean number of crossovers per meiosis (e.g. size in Morgan of the chromosome)
markers	If a single number is provided, the number is used as the total number of markers generated either randomly, or using a regular distribution (a regular distribution is chosen if the number is negative). If a vector is provided, that vector is used.

time_points	vector with time points at which local ancestry has to be recorded to be returned at the end of the simulation. If left at -1, ancestry is recorded at every generation (computationally heavy).
num_threads	default is 1. -1 takes all available threads.
verbose	displays a progress bar
record_true_junctions	keep track of the true number of junctions?
num_indiv_sampled	the number of individuals sampled at each time point to be genotyped
coverage	fraction of markers that can be successfully phased
error_rate	fraction of markers that are erroneously phased (e.g. swapped)

Value

a tibble with five columns: [time, individual, marker location, ancestry chromosome 1, ancestry chromosome 2]

Examples

```
## Not run:
sim_phased_unphased(pop_size = 100, freq_ancestor_1 = 0.5,
                    total_runtime = 10, size_in_morgan = 1,
                    markers = 10, time_points = c(0, 5, 10),
                    num_threads = 1)

## End(Not run)
```

time_error	<i>Estimate the error in the time estimate</i>
------------	--

Description

Calculate the error in the estimate of the onset of hybridization, following Equations 3 & 4 in the Supplementary information of Janzen et al. 2018.

Usage

```
time_error(t = NA, N = Inf, R = Inf, H_0 = 0.5, C = 1, relative = TRUE)
```

Arguments

t	Inferred time
N	Population Size
R	Number of genetic markers
H_0	Frequency of heterozygosity at t = 0

C	Mean number of crossovers per meiosis (e.g. size in Morgan of the chromosome)
relative	Boolean flag, if TRUE: return the relative error, if FALSE: return error in generations

Value

Expected error in the time estimate

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