# **Package: simRestore (via r-universe)**

August 30, 2024

Type Package

Title Simulate the Effect of Management Policies on Restoration Efforts

Version 1.1.4

**Description** Simulation methods to study the effect of management policies on efforts to restore populations back to their original genetic composition. Allows for single-scenario simulation and for optimization of specific chosen scenarios. Further information can be found in Hernandez, Janzen and Lavretsky (2023) <doi:10.1111/1755-0998.13892>.

**License** GPL ( $\geq 2$ )

Imports Rcpp, shiny, subplex, tibble

**Suggests** dplyr, ggplot2, knitr, magrittr, rmarkdown, shinyBS, shinythemes, shinyWidgets, testthat, tidyr, egg

LinkingTo Rcpp

VignetteBuilder knitr

Encoding UTF-8

RoxygenNote 7.2.3

Repository https://thijsjanzen.r-universe.dev

RemoteUrl https://github.com/thijsjanzen/simrestore

RemoteRef HEAD

RemoteSha 0176a5d0942bdf4a952b4fe648fe10bd241e3e88

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optimize\_adaptive

Optimize a policy assuming a fixed total sum across all generations of individuals that can be put or pulled (e.g. a fixed effort). This fixed total sum is distributed across the generations following a beta distribution, and the parameters of this beta distribution are fitted.

# Description

Optimize a policy assuming a fixed total sum across all generations of individuals that can be put or pulled (e.g. a fixed effort). This fixed total sum is distributed across the generations following a beta distribution, and the parameters of this beta distribution are fitted.

# Usage

```
optimize_adaptive(
  target_frequency = 0.99,
  initial_population_size = 400,
  reproduction_success_rate = 0.387,
  reproductive_risk = c(0.2, 0),
  K = 400,
  num_generations = 20,
  optimize_put = 100,
  optimize_pull = 0,
  starting_freq = 0.2,
  sd_starting_freq = 0.05,
  morgan = c(1),
  establishment_burnin = 30,
  num_replicates = 1,
  max_age = 6,
  mean_number_of_offspring = 6,
  sd_number_of_offspring = 1,
  genetic_model = "point",
  smin = 0.5,
  smax = 0.9,
  b = -2,
  p = 0.5,
  sex_ratio_put = 0.5,
  sex_ratio_pull = 0.5,
  sex_ratio_offspring = 0.5,
  ancestry_put = 1,
  ancestry_pull = 1,
  random_mating = FALSE,
  extra_pair_copulation = 0,
  verbose = FALSE,
  return_genetics = FALSE
)
```

# Arguments

target\_frequency

frequency to aim for

# initial\_population\_size

population size at the start

# reproduction\_success\_rate

reproduction_success_rate			
	frequency of females that yield offspring at the end of the breeding season (e.g. a fraction of 1 - reproduction_success_rate of females. This is a joint effect of breeding females getting killed (see female_death_rate) and other sources of failure to produce offspring. Other sources of failure are calculated from reproduction_success_rate and female_death_rate, such that the resulting reproduction failure rate = 1 - reproduction_success_rate / (1 - female breeding risk)		
reproductive_ri	sk		
	Additional death rate of males and females as a result of breeding (e.g. as a result of protecting the offspring against predators). Provide as a vector where the first index indicates the risk for females, the second the risk for males.		
К	carrying capacity		
num_generations			
	number of generations		
optimize_put	optimization proceeds such that the sum of all addition over all generations is equal to this number. Switch off by setting to zero. The individuals are dis- tributed over time following a beta distribution.		
optimize_pull	Optimization proceeds such that the sum of all removal is equal to this number. Switch off by setting to zero.		
starting_freq	initial frequency of target ancestry in the population.		
sd_starting_fre	pq		
	variation in initial frequency of target ancestry.		
morgan	a vector with the size of each chromosome in morgan, e.g. if a single chromosome is to be simulated a single number will suffice, but for two chromosomes of a size of 1 Morgan, a vector like: $c(1, 1)$ will work.		
establishment_b	burnin		
	number of generations before establishment		
num_replicates	number of replicates per parameter combination to be simulated. Fit of the parameter combination is chosen as the average frequency across replicates.		
max_age	maximum age an individual can reach.		
<pre>mean_number_of_</pre>	offspring		
	mean number of offspring per female		
<pre>sd_number_of_of</pre>	fspring		
	standard deviation of number of offspring per female (assuming the number of offspring is always 0 or larger).		
genetic_model	The model can either use the point ancestry model ("point") of underlying genet- ics, which speeds up simulation considerably, but underestimates genetic varia- tion. Alternatively, a more detailed genetic model is available, making use of the theory of junctions, this can be accessed using the option "junctions". Default is "point".		

smin	minimum survival rate	
smax	maximum survival rate	
b	steepness of the survival rate. Negative values indicate a declining survival rate with increasing population size, positive values indicate an increasing survival rate with increasing population size.	
р	Density at which the survival rate changes most relative. Expressed in N / K (e.g., for a value of 1.0, the survival rate changes most rapidly around N = K, for a value of 0.5, the survival rate changes most rapidly around N = 0.5K, etc).	
sex_ratio_put	sex ratio of individuals that are added (if any) to the population. Sex ratio is expressed as males / (males + females), such that 0.5 indicates an even sex ratio, 0.9 indicates a male biased sex ratio and 0.1 indicates a female biased sex ratio.	
sex_ratio_pull	sex ratio of individuals that are removed (if any) from the population. The sex ratio is expressed as males / (males + females), such that 0.5 indicates an even sex ratio, 0.9 indicates a male biased sex ratio and 0.1 indicates a female biased sex ratio.	
<pre>sex_ratio_offsp</pre>	pring	
	sex ratio of newly born offspring. The sex ratio is expressed as males / (males + females), such that 0.5 indicates an even sex ratio, 0.9 indicates a male biased sex ratio and 0.1 indicates a female biased sex ratio.	
ancestry_put	Average ancestry of individuals being used for supplementation. If the target is high focal ancestry (e.g. aiming for focal ancestry of 1.0), ancestry put should reflect this and be set to 1.0 ( which is the default value). When supplementing with non-pure individuals, this value can consequently be lowered.	
ancestry_pull	Ancestry level below which individuals are allowed to be pulled - this can reduce the effective number of individuals pulled if none of the individuals in the pop- ulation match this ancestry level. This can be used to selectively only remove those with low target ancestry.	
random_mating	by default, simulations assume fixed pair bonding, e.g. each female mates with exactly one male (if available). Alternatively, if random_mating = TRUE, females will mate with a random male, introducing the possibility that some males mate multiple times.	
extra_pair_copu	lation	
	probability of offspring to be fathered by another male. We assume that all offspring from one mother can have at most two fathers.	
verbose	provides verbose output if TRUE.	
return_genetics		
	returns a tibble containing all local ancestry information for all individuals. This tibble contains the following informative columns: 1) time (the last generation), 2) replicate, 3) individual, 4) sex ( $0 = male$ , $1 = female$ ), 5) Linkage Group (if use_simplified_model == FALSE), 6) chromosome (1 or 2, returning phased results), 7) position (if use_simplified_model == FALSE) and 8) local ancestry (0 or 1).	

# Value

list with five elements: 1) put: optimal number of individuals to put (0 if not estimated), 2) pull: optimal number of individuals to pull (0 if not estimated), 3) results tibble (see simulate\_policy()),

# optimize\_static

4) curve tibble with three columns, indicating the realized number of put/pull per generation, with column 1) time in generations, column 2) number of individuals to put in generation t and 3) number of individuals to pull in generation t. The last element of the list contains the final obtained frequency for the best fit.

## Examples

```
opt_res <- optimize_adaptive(target_frequency = 0.99,</pre>
                              optimize_put = 1000,
                              num_generations = 20,
                              starting_freq = 0.2,
                              initial_population_size = 100)
opt_res$put
```

optimize\_static

Optimize putting and/or pulling, where it is assumed that the same amount is applied per generation.

# Description

Optimize putting and/or pulling, where it is assumed that the same amount is applied per generation.

# Usage

```
optimize_static(
  target_frequency = 0.99,
  initial_population_size = 400,
  reproduction_success_rate = 0.387,
  reproductive_risk = c(0.2, 0),
 K = 400,
  num_generations = 20,
 optimize_put = TRUE,
 optimize_pull = FALSE,
  starting_freq = 0.2,
  sd_starting_freq = 0.05,
 morgan = c(1),
  establishment_burnin = 30,
  num_replicates = 1,
 max_age = 6,
 mean_number_of_offspring = 6,
  sd_number_of_offspring = 1,
  genetic_model = "point",
  smin = 0.5,
  smax = 0.9,
  b = -2,
  p = 0.5,
  sex_ratio_put = 0.5,
  sex_ratio_pull = 0.5,
```

```
sex_ratio_offspring = 0.5,
  ancestry_put = 1,
  ancestry_pull = 1,
  random_mating = FALSE,
  extra_pair_copulation = 0,
  verbose = FALSE,
  return_genetics = FALSE
)
```

Arguments target\_frequency frequency to aim for initial\_population\_size population size at the start reproduction\_success\_rate frequency of females that yield offspring at the end of the breeding season (e.g. a fraction of 1 - reproduction\_success\_rate of females. This is a joint effect of breeding females getting killed (see female\_death\_rate) and other sources of failure to produce offspring. Other sources of failure are calculated from reproduction success rate and female death rate, such that the resulting reproduction failure rate = 1 - reproduction\_success\_rate / (1 - female breeding risk) reproductive\_risk Additional death rate of males and females as a result of breeding (e.g. as a result of protecting the offspring against predators). Provide as a vector where the first index indicates the risk for females, the second the risk for males. Κ carrying capacity num\_generations number of generations When set to 0, FALSE or a negative number, it will not be optimized. When negoptimize\_put ative, the absolute value will be taken as a fixed contribution to each generation (but will not be optimized) When set to 0, FALSE or a negative number, it will not be optimized. When negoptimize\_pull ative, the absolute value will be taken as a fixed contribution to each generation (but will not be optimized) starting\_freq initial frequency of target ancestry in the population. sd\_starting\_freq variation in initial frequency of target ancestry. a vector with the size of each chromosome in morgan, e.g. if a single chromomorgan some is to be simulated a single number will suffice, but for two chromosomes of a size of 1 Morgan, a vector like: c(1, 1) will work. establishment\_burnin number of generations before establishment num\_replicates number of replicates maximum age an individual can reach. max\_age

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# optimize\_static

<pre>mean_number_of_</pre>	offspring	
	mean number of offspring per female	
sd_number_of_of	fspring standard deviation of number of offspring per female (assuming the number of offspring is always 0 or larger).	
genetic_model	The model can either use the point ancestry model ("point") of underlying genet- ics, which speeds up simulation considerably, but underestimates genetic varia- tion. Alternatively, a more detailed genetic model is available, making use of the theory of junctions, this can be accessed using the option "junctions". Default is "point".	
smin	minimum survival rate	
smax	maximum survival rate	
b	steepness of the survival rate. Negative values indicate a declining survival rate with increasing population size, positive values indicate an increasing survival rate with increasing population size.	
р	Density at which the survival rate changes most relative. Expressed in N / K (e.g., for a value of 1.0, the survival rate changes most rapidly around N = K, for a value of 0.5, the survival rate changes most rapidly around N = 0.5K, etc).	
sex_ratio_put	sex ratio of individuals that are added (if any) to the population. Sex ratio is expressed as males / (males + females), such that 0.5 indicates an even sex ratio, 0.9 indicates a male biased sex ratio and 0.1 indicates a female biased sex ratio.	
sex_ratio_pull	sex ratio of individuals that are removed (if any) from the population. The sex ratio is expressed as males / (males + females), such that 0.5 indicates an even sex ratio, 0.9 indicates a male biased sex ratio and 0.1 indicates a female biased sex ratio.	
<pre>sex_ratio_offsp</pre>	pring	
	sex ratio of newly born offspring. The sex ratio is expressed as males / (males + females), such that 0.5 indicates an even sex ratio, 0.9 indicates a male biased sex ratio and 0.1 indicates a female biased sex ratio.	
ancestry_put	Average ancestry of individuals being used for supplementation. If the target is high focal ancestry (e.g. aiming for focal ancestry of 1.0), ancestry put should reflect this and be set to 1.0 ( which is the default value). When supplementing with non-pure individuals, this value can consequently be lowered.	
ancestry_pull	Ancestry level below which individuals are allowed to be pulled - this can reduce the effective number of individuals pulled if none of the individuals in the pop- ulation match this ancestry level. This can be used to selectively only remove those with low target ancestry.	
random_mating	by default, simulations assume fixed pair bonding, e.g. each female mates with exactly one male (if available). Alternatively, if random_mating = TRUE, females will mate with a random male, introducing the possibility that some males mate multiple times.	
extra_pair_copulation		
	probability of offspring to be fathered by another male. We assume that all offspring from one mother can have at most two fathers.	
verbose	provides verbose output if TRUE.	

### return\_genetics

returns a tibble containing all local ancestry information for all individuals. This tibble contains the following informative columns: 1) time (the last generation), 2) replicate, 3) individual, 4) sex (0 = male, 1 = female), 5) Linkage Group (if use simplified model == FALSE), 6) chromosome (1 or 2, returning phased results), 7) position (if use\_simplified\_model == FALSE) and 8) local ancestry (0 or 1).

# Value

list with five elements: 1) put: optimal number of individuals to put (0 if not estimated), 2) pull: optimal number of individuals to pull (0 if not estimated), 3) results tibble (see simulate\_policy()), 4) curve tibble with three columns, indicating the realized number of put/pull per generation, with column 1) time in generations, column 2) number of individuals to put in generation t and 3) number of individuals to pull in generation t. The last element of the list contains the final obtained frequency for the best fit.

# Examples

```
opt_res <- optimize_static(target_frequency = 0.99,</pre>
                            optimize_put = TRUE,
                            num_generations = 10,
                            starting_freq = 0.2,
                            initial_population_size = 100)
```

opt\_res\$put

run\_shiny\_app runs shiny app locally

# Description

This function allows for local execution of the shiny app. Alternatively, an online version of this app can be found here.

# Usage

run\_shiny\_app()

# Value

No return value

simulate\_policy

# Description

Using this function, the user can simulate the effect of an intended management policy on the genetic composition of a focal population. The population is assumed to have overlapping generations, and the user can specify two genetic models, either using a simplified average ancestry representation (genetic\_model = "point"), or a more detailed model tracking explicit recombination among chromosomes, using genetic\_model = "junctions".

# Usage

```
simulate_policy(
  initial_population_size = 400,
  reproduction_success_rate = 0.387,
  reproductive_risk = c(0.2, 0),
 K = 400,
  num_generations = 20,
  pull = 0,
  put = 0,
  starting_freq = 0.5,
  sd_starting_freq = 0.05,
 morgan = c(1),
 max_age = 6,
 mean_number_of_offspring = 6,
  sd_number_of_offspring = 1,
  genetic_model = "point",
  establishment_burnin = 30,
  num_replicates = 1,
  seed = NULL,
  smin = 0.5,
  smax = 0.9,
 b = -2,
  p = 0.5,
  sex_ratio_put = 0.5,
  sex_ratio_pull = 0.5,
  sex_ratio_offspring = 0.5,
  ancestry_put = 1,
  ancestry_pull = 1,
  random_mating = FALSE,
  extra_pair_copulation = 0,
  verbose = FALSE,
  return_genetics = FALSE
)
```

# Arguments

initial_populat	ion_size
	population size at the start
reproduction_su	iccess_rate
	frequency of females that yield offspring at the end of the breeding season (e.g. a fraction of 1 - reproduction_success_rate of females. This is a joint effect of breeding females getting killed (see female_death_rate) and other sources of failure to produce offspring. Other sources of failure are calculated from reproduction_success_rate and female_death_rate, such that the resulting reproduction failure rate = 1 - reproduction_success_rate / (1 - female breeding risk)
reproductive_ri	sk
	Additional death rate of males and females as a result of breeding (e.g. as a result of protecting the offspring against predators). Provide as a vector where the first index indicates the risk for females, the second the risk for males.
К	carrying capacity
num_generations	i
	number of generations
pull	vector of the number of individuals pulled per year
put	vector of the number of individuals added per year
<pre>starting_freq sd_starting_fre</pre>	initial frequency of target ancestry in the population.
5-	variation in initial frequency of target ancestry.
morgan	a vector with the size of each chromosome in morgan, e.g. if a single chromosome is to be simulated a single number will suffice, but for two chromosomes of a size of 1 Morgan, a vector like: $c(1, 1)$ will work.
max_age	maximum age an individual can reach.
<pre>mean_number_of_</pre>	offspring
	mean number of offspring per female
<pre>sd_number_of_of</pre>	fspring
	standard deviation of number of offspring per female (assuming the number of offspring is always 0 or larger).
genetic_model	The model can either use the point ancestry model ("point") of underlying genet- ics, which speeds up simulation considerably, but underestimates genetic varia- tion. Alternatively, a more detailed genetic model is available, making use of the theory of junctions, this can be accessed using the option "junctions". Default is "point".
establishment_b	purnin
	number of generations before establishment
num_replicates	number of replicates
seed	random number seed, if left open, current time is used.
smin	minimum survival rate
smax	maximum survival rate

b	steepness of the survival rate. Negative values indicate a declining survival rate with increasing population size, positive values indicate an increasing survival rate with increasing population size.
p	Density at which the survival rate changes most relative. Expressed in N / K (e.g., for a value of 1.0, the survival rate changes most rapidly around N = K, for a value of 0.5, the survival rate changes most rapidly around N = 0.5K, etc).
sex_ratio_put	sex ratio of individuals that are added (if any) to the population. Sex ratio is expressed as males / (males + females), such that 0.5 indicates an even sex ratio, 0.9 indicates a male biased sex ratio and 0.1 indicates a female biased sex ratio.
<pre>sex_ratio_pull</pre>	sex ratio of individuals that are removed (if any) from the population. The sex ratio is expressed as males / (males + females), such that 0.5 indicates an even sex ratio, 0.9 indicates a male biased sex ratio and 0.1 indicates a female biased sex ratio.
<pre>sex_ratio_offsp</pre>	pring
	sex ratio of newly born offspring. The sex ratio is expressed as males / (males + females), such that 0.5 indicates an even sex ratio, 0.9 indicates a male biased sex ratio and 0.1 indicates a female biased sex ratio.
ancestry_put	Average ancestry of individuals being used for supplementation. If the target is high focal ancestry (e.g. aiming for focal ancestry of 1.0), ancestry put should reflect this and be set to 1.0 ( which is the default value). When supplementing with non-pure individuals, this value can consequently be lowered.
ancestry_pull	Ancestry level below which individuals are allowed to be pulled - this can reduce the effective number of individuals pulled if none of the individuals in the pop- ulation match this ancestry level. This can be used to selectively only remove those with low target ancestry.
random_mating	by default, simulations assume fixed pair bonding, e.g. each female mates with exactly one male (if available). Alternatively, if random_mating = TRUE, females will mate with a random male, introducing the possibility that some males mate multiple times.
extra_pair_copu	ulation
	probability of offspring to be fathered by another male. We assume that all offspring from one mother can have at most two fathers.
verbose	provides verbose output if TRUE.
return_genetics	3
-	returns a tibble containing all local ancestry information for all individuals. This tibble contains the following informative columns: 1) time (the last generation), 2) replicate, 3) individual, 4) sex (0 = male, 1 = female), 5) Linkage Group (if use_simplified_model == FALSE), 6) chromosome (1 or 2, returning phased results), 7) position (if use_simplified_model == FALSE) and 8) local ancestry (0 or 1).

# Value

tibble with 8 columns: 1) replicate, 2) time (in generations), 3) average frequency of ancestry across all individuals 4) average frequency of ancestry across all males, 5) average frequency of ancestry across all females, 6) number of individuals, 7) number of males and 8) number of females if

return\_genetics = TRUE, the output is a list containing the above mentioned tibble, called 'results', and a second tibble called 'genetics', with the local ancestry in long format, split out per generation, replicate, individual, sex, linkage group and chromosome (1 or 2). Here, linkage group indicates the focal chromosome (linkage group), and 'chromosome' indicates which of the diploid pair of chromosomes is measured, allowing for phased output if required.

# Examples

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