

# Package ‘fwsim’

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**Version** 0.3.4

**Title** Fisher-Wright Population Simulation

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**Description** Simulates a population under the Fisher-Wright model (fixed or stochastic population size) with a one-step neutral mutation process (stepwise mutation model, logistic mutation model and exponential mutation model supported). The stochastic population sizes are random Poisson distributed and different kinds of population growth are supported. For the stepwise mutation model, it is possible to specify locus and direction specific mutation rate (in terms of upwards and downwards mutation rate). Intermediate generations can be saved in order to study e.g. drift.

**License** GPL-2

**Depends** R (>= 3.1.0)

**LinkingTo** Rcpp

**Imports** Rcpp (>= 0.11), methods

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 fwsim-package

*Haplotype tools*


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### Description

Tools for analysing haplotypes, e.g. population simulation.

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 fwsim

*Fisher-Wright Population Simulation*


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### Description

This package provides tools to simulate a population under the Fisher-Wright model with a stepwise neutral mutation process on  $r$  loci, where mutations on loci happen independently. The population sizes are either fixed (traditional/original Fisher-Wright model) or random Poisson distributed with exponential growth supported. Intermediate generations can be saved in order to study e.g. drift.

For stochastic population sizes: Model described in detail at <http://arxiv.org/abs/1210.1773>. Let  $M$  be the population size at generation  $i$  and  $N$  the population size at generation  $i + 1$ . Then we assume that  $N$  conditionally on  $M$  is Poisson( $\alpha M$ ) distributed for  $\alpha > 0$  ( $\alpha > 1$  gives expected growth and  $0 < \alpha < 1$  gives expected decrease).

For each haplotype  $x$  occurring  $m$  times in the  $i$ 'th generation, the number of children  $n$  is Poisson( $\alpha m$ ) distributed independently of other haplotypes. It then follows that the sum of the number of haplotypes follows a Poisson( $\alpha M$ ) distribution (as just stated in the previous paragraph) and that  $n$  conditionally on  $N$  follows a Binomial( $N, m/M$ ) as expected.

The mutation model can be e.g. the stepwise neutral mutation model. See `init_mutmodel` for details.

### Usage

```
fwsim(G, H0, N0, mutmodel, alpha = 1.0, SNP = FALSE,
      save_generations = NULL, progress = TRUE, trace = FALSE, ensure_children = FALSE, ...)
fwsim_fixed(G, H0, N0, mutmodel, SNP = FALSE,
            save_generations = NULL, progress = TRUE, trace = FALSE, ...)
## S3 method for class 'fwsim'
print(x, ...)
## S3 method for class 'fwsim'
summary(object, ...)
## S3 method for class 'fwsim'
plot(x, which = 1L, ...)
```

**Arguments**

G	number of generations to evolve (integer, remember postfix L).
H0	haplotypes of the initial population. Must be a vector or matrix (if more than one initial haplotype). The number of loci is the length or number of columns of H0.
N0	count of the H0 haplotypes. The i'th element is the count of the haplotype H0[i, ]. $\text{sum}(N0)$ is the size of initial population.
mutmodel	a mutmodel object created with <code>init_mutmodel</code> . Alternatively, a numeric vector of length r of mutation probabilities (this will create a stepwise mutation model with r loci divide the mutation probabilities evenly between upwards and downwards mutation).
alpha	vector of length 1 or G of growth factors (1 correspond to expected constant population size). If length 1, the value is reused in creating a vector of length G.
SNP	to make alleles modulus 2 to immitate SNPs.
save_generations	to save intermediate populations. NULL means that no intermediate population will be saved. Else, a vector of the generation numbers to save.
progress	whether to print progress of the evolution.
trace	whether to print trace of the evolution (more verbose than progress).
ensure_children	Ensures that every generation has at least one child; implemented by getting $\text{Poisson}(\alpha M) + 1$ children.
x	A fwsim object.
object	A fwsim object.
which	A number specifying the plot (currently only 1: the actual population sizes vs the expected sizes).
...	not used.

**Value**

A fwsim object with elements

pars	the parameters used for the simulation
saved_populations	a list of haplotypes in the intermediate populations
population	haplotypes in the end population after G generations
pop_sizes	the population size for each generation
expected_pop_sizes	the expected population size for each generation

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**Examples**

```

# SMM (stepwise mutation model) example
set.seed(1)
fit <- fwsim(G = 100L, H0 = c(0L, 0L, 0L), N0 = 10000L,
  mutmodel = c(Loc1 = 0.001, Loc2 = 0.002, Loc3 = 0.003))
summary(fit)
fit

# SMM (stepwise mutation model) example
H0 <- matrix(c(0L, 0L, 0L), 1L, 3L, byrow = TRUE)

mutmodel <- init_mutmodel(modeltype = 1L,
  mutpars = matrix(c(c(0.003, 0.001), rep(0.004, 2), rep(0.001, 2)),
    ncol = 3,
    dimnames = list(NULL, c("DYS19", "DYS389I", "DYS391"))))
mutmodel

set.seed(1)
fit <- fwsim(G = 100L, H0 = H0, N0 = 10000L, mutmodel = mutmodel)

xtabs(N ~ DHS19 + DHS389I, fit$population)
plot(1L:fit$pars$G, fit$pop_sizes, type = "l",
  ylim = range(range(fit$pop_sizes), range(fit$expected_pop_sizes)))
points(1L:fit$pars$G, fit$expected_pop_sizes, type = "l", col = "red")

set.seed(1)
fit_fixed <- fwsim_fixed(G = 100L, H0 = H0, N0 = 10000L, mutmodel = mutmodel)

# LMM (logistic mutation model) example
mutpars.locus1 <- c(0.149, 2.08, 18.3, 0.149, 0.374, 27.4) # DHS19
mutpars.locus2 <- c(0.500, 1.18, 18.0, 0.500, 0.0183, 349) # DHS389I
mutpars.locus3 <- c(0.0163, 17.7, 11.1, 0.0163, 0.592, 14.1) # DHS391
mutpars <- matrix(c(mutpars.locus1, mutpars.locus2, mutpars.locus3), ncol = 3)
colnames(mutpars) <- c("DYS19", "DYS389I", "DYS391")
mutmodel <- init_mutmodel(modeltype = 2L, mutpars = mutpars)
mutmodel

set.seed(1)
H0_LMM <- matrix(c(15L, 13L, 10L), 1L, 3L, byrow = TRUE)
fit_LMM <- fwsim(G = 100L, H0 = H0_LMM, N0 = 10000L, mutmodel = mutmodel)
xtabs(N ~ DHS19 + DHS389I, fit_LMM$population)

```

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init\_mutmodel

*init\_mutmodel*


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**Description**

Method to initialise a mutation model.

**Usage**

```
init_mutmodel(modeltype = 1, mutpars = NULL, ...)
## S3 method for class 'mutmodel'
print(x, ...)
```

**Arguments**

modeltype	1: SMM (traditional single-step mutation model). 2: LMM (Logistic mutation model introduced in Jochens (2011) 'Empirical Evaluation Reveals Best Fit of a Logistic Mutation Model for Human Y-Chromosomal Microsatellites'). 3: Exponential mutation model (unpublished).
mutpars	A matrix specifying the mutation parameters for each locus. Rows are parameters and columns are loci. If a vector, the same values are used for all loci.
x	A mutmodel object.
...	not used.

**Details**

Mutation parameters for each locus.

Mutmodel 1 (SMM): 2 parameters per locus

$$P(i \rightarrow i-1) = \mu\_d$$

$$P(i \rightarrow i+1) = \mu\_u$$

$$P(i \rightarrow i) = 1 - P(i \rightarrow i-1) - P(i \rightarrow i+1)$$

$$= 1 - \mu\_d - \mu\_u$$

mutpars[1, locus]:  $\mu\_d$   
mutpars[2, locus]:  $\mu\_u$

Mutmodel 2 (LMM): 6 parameters per locus

$$P(i \rightarrow i-1) = \gamma\_d / (1 + \exp(\alpha\_d * (\beta\_d - i)))$$

$$P(i \rightarrow i+1) = \gamma\_u / (1 + \exp(\alpha\_u * (\beta\_u - i)))$$

$$P(i \rightarrow i) = 1 - P(i \rightarrow i-1) - P(i \rightarrow i+1)$$

mutpars[1, locus]:  $\gamma\_d$   
mutpars[2, locus]:  $\alpha\_d$   
mutpars[3, locus]:  $\beta\_d$   
mutpars[4, locus]:  $\gamma\_u$   
mutpars[5, locus]:  $\alpha\_u$   
mutpars[6, locus]:  $\beta\_u$

Mutmodel 3 (EMM): 4 parameters per locus

$$P(i \rightarrow i-1) = 1 / ((1 + \exp(a + b*i)) * (1 + \exp(\alpha + \beta*i)))$$

$$P(i \rightarrow i+1) = \exp(\alpha + \beta*i) / ((1 + \exp(a + b*i)) * (1 + \exp(\alpha + \beta*i)))$$

$$P(i \rightarrow i) = 1 - P(i \rightarrow i-1) - P(i \rightarrow i+1)$$

$$= \exp(a + b*i) / (1 + \exp(a + b*i))$$

```
mutpars[1, locus]: a
mutpars[2, locus]: b
mutpars[3, locus]: alpha
mutpars[4, locus]: beta
```

### Value

A mutmodel object (a list with entire model type and mutpars).

### Examples

```
mutpars <- matrix(c(c(0.003, 0.001), rep(0.004, 2), rep(0.001, 2)), ncol = 3)
colnames(mutpars) <- c("DYS19", "DYS389I", "DYS391")
mutmodel <- init_mutmodel(modeltype = 1L, mutpars = mutpars)
mutmodel

mutpars.locus1 <- c(0.149, 2.08, 18.3, 0.149, 0.374, 27.4) # DYS19
mutpars.locus2 <- c(0.500, 1.18, 18.0, 0.500, 0.0183, 349) # DYS389I
mutpars.locus3 <- c(0.0163, 17.7, 11.1, 0.0163, 0.592, 14.1) # DYS391
mutpars <- matrix(c(mutpars.locus1, mutpars.locus2, mutpars.locus3), ncol = 3)
colnames(mutpars) <- c("DYS19", "DYS389I", "DYS391")
mutmodel <- init_mutmodel(modeltype = 2L, mutpars = mutpars)
mutmodel
```

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mutmodel	<i>Mutation model logic</i>
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### Description

Functions for mutation model logic, e.g. probability of downwards and upwards mutations etc.

### Usage

```
mutmodel_not_mut(mutmodel, locus, alleles)
mutmodel_dw_mut(mutmodel, locus, alleles)
mutmodel_up_mut(mutmodel, locus, alleles)
approx_stationary_dist(mutmodel, alleles)
```

### Arguments

mutmodel	a mutmodel object created with <code>init_mutmodel</code> .
locus	the locus of interest (integer, remember postfix L).
alleles	vector of integers (remember postfix L) of the alleles of interest.

### Value

Mutation probabilities for locus locus in mutation model mutmodel at alleles alleles.

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**Examples**

```
mutpars.locus1 <- c(0.149, 2.08, 18.3, 0.149, 0.374, 27.4) # DYS19
mutpars.locus2 <- c(0.500, 1.18, 18.0, 0.500, 0.0183, 349) # DYS389I
mutpars.locus3 <- c(0.0163, 17.7, 11.1, 0.0163, 0.592, 14.1) # DYS391
mutpars <- matrix(c(mutpars.locus1, mutpars.locus2, mutpars.locus3), ncol = 3)
colnames(mutpars) <- c("DYS19", "DYS389I", "DYS391")
mutmodel <- init_mutmodel(modeltype = 2L, mutpars = mutpars)
```

```
mutmodel_not_mut(mutmodel, locus = 1L, alleles = 10L:20L)
mutmodel_dw_mut(mutmodel, locus = 1L, alleles = 10L:20L)
mutmodel_up_mut(mutmodel, locus = 1L, alleles = 10L:20L)
```

```
statdists <- approx_stationary_dist(mutmodel, alleles = 5L:20L)
bp <- barplot(statdists, beside = TRUE)
text(bp, 0.02, round(statdists, 1), cex = 1, pos = 3)
text(bp, 0, rep(rownames(statdists), ncol(mutmodel$mutpars)), cex = 1, pos = 3)
```

```
mutpars <- matrix(c(c(0.003, 0.001), rep(0.004, 2), rep(0.001, 2)), ncol = 3)
colnames(mutpars) <- c("DYS19", "DYS389I", "DYS391")
mutmodel <- init_mutmodel(modeltype = 1L, mutpars = mutpars)
mutmodel
statdists <- approx_stationary_dist(mutmodel, alleles = 5L:20L)
statdists
```

```
bp <- barplot(statdists, beside = TRUE)
text(bp, 0.02, round(statdists, 1), cex = 1, pos = 3)
text(bp, 0, rep(rownames(statdists), ncol(mutmodel$mutpars)), cex = 1, pos = 3)
```

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