

# Package ‘OncoSimulR’

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**Type** Package

**Title** Forward Genetic Simulation of Cancer Progression with Epistasis

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**Description** Functions for forward population genetic simulation in asexual populations, with special focus on cancer progression. Fitness can be an arbitrary function of genetic interactions between multiple genes or modules of genes, including epistasis, order restrictions in mutation accumulation, and order effects. Mutation rates can differ between genes, and we can include mutator/antimutator genes (to model mutator phenotypes). Simulations use continuous-time models and can include driver and passenger genes and modules. Also included are functions for: simulating random DAGs of the type found in Oncogenetic Tress, Conjunctive Bayesian Networks, and other tumor progression models; plotting and sampling from single or multiple realizations of the simulations, including single-cell sampling; plotting the parent-child relationships of the clones; generating random fitness landscapes (Rough Mount Fuji, House of Cards, and additive models) and plotting them.

**biocViews** BiologicalQuestion, SomaticMutation

**License** GPL (>= 3)

**URL** <https://github.com/rdiaz02/OncoSimul>,  
<https://popmodels.cancercontrol.cancer.gov/gsr/packages/oncosimulr/>

**BugReports** <https://github.com/rdiaz02/OncoSimul/issues>

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allFitnessEffects	<i>Create fitness and mutation effects specification from restrictions, epistasis, and order effects.</i>
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### Description

Given one or more of a set of poset restrictions, epistatic interactions, order effects, and genes without interactions, as well as, optionally, a mapping of genes to modules, return the complete fitness specification.

For mutator effects, given one or more of a set of epistatic interactions and genes without interactions, as well as, optionally, a mapping of genes to modules, return the complete specification of how mutations affect the mutation rate.

The output of these functions is not intended for user consumption, but as a way of preparing data to be sent to the C++ code.

## Usage

```
allFitnessEffects(rT = NULL, epistasis = NULL, orderEffects = NULL,
  noIntGenes = NULL, geneToModule = NULL, drvNames = NULL,
  genotFitness = NULL, keepInput = TRUE)

allMutatorEffects(epistasis = NULL, noIntGenes = NULL,
  geneToModule = NULL,
  keepInput = TRUE)
```

## Arguments

**rT** A restriction table that is an extended version of a poset (see [poset](#)). A restriction table is a data frame where each row shows one edge between a parent and a child. A restriction table contains exactly these columns, in this order:

- parent** The identifiers of the parent nodes, in a parent-child relationship. There must be at least one entry with the name "Root".
- child** The identifiers of the child nodes.
- s** A numeric vector with the fitness effect that applies if the relationship is satisfied.
- sh** A numeric vector with the fitness effect that applies if the relationship is not satisfied. This provides a way of explicitly modeling deviations from the restrictions in the graph, and is discussed in Diaz-Uriarte, 2015.
- typeDep** The type of dependency. Three possible types of relationship exist:
  - AND, monotonic, or CMPN** Like in the CBN model, all parent nodes must be present for a relationship to be satisfied. Specify it as "AND" or "MN" or "monotone".
  - OR, semimonotonic, or DMPN** A single parent node is enough for a relationship to be satisfied. Specify it as "OR" or "SM" or "semimonotone".
  - XOR or XMPN** Exactly one parent node must be mutated for a relationship to be satisfied. Specify it as "XOR" or "xmpn" or "XMPN".
 In addition, for the nodes that depend only on the root node, you can use "-" or "-" if you want (though using any of the other three would have the same effects if a node that connects to root only connects to root).

**epistasis** A named numeric vector. The names identify the relationship, and the numeric value is the fitness (or mutator) effect. For the names, each of the genes or modules involved is separated by a ":". A negative sign denotes the absence of that term.

**orderEffects** A named numeric vector, as for epistasis. A ">" separates the names of the genes or modules of a relationship, so that "U > Z" means that the relationship is satisfied when mutation U has happened before mutation Z.

**noIntGenes** A numeric vector (optionally named) with the fitness coefficients (or mutator multiplier factor) of genes (only genes, not modules) that show no interactions. These genes cannot be part of modules. But you can specify modules that have no epistatic interactions. See examples and vignette.

Of course, avoid using potentially confusing characters in the names. In particular, ",", " and ">" are not allowed as gene names.

**geneToModule** A named character vector that allows to match genes and modules. The names are the modules, and each of the values is a character vector with the gene names,

separated by a comma, that correspond to a module. Note that modules cannot share genes. There is no need for modules to contain more than one gene. If you specify a `geneToModule` argument, and you used a restriction table, the `geneToModule` must necessarily contain, in the first position, "Root" (since the restriction table contains a node named "Root"). See examples below.

<code>drvNames</code>	The names of genes that are considered drivers. This is only used for: a) deciding when to stop the simulations, in case you use number of drivers as a simulation stopping criterion (see <code>oncoSimulIndiv</code> ); b) for summarization purposes (e.g., how many drivers are mutated); c) in figures. But you need not specify anything if you do not want to, and you can pass an empty vector (as <code>character(0)</code> ). The default has changed with respect to v.2.1.3 and previous: it used to be to assume that all genes that were not in the <code>noIntGenes</code> were drivers. The default now is to assume nothing: if you want <code>drvNames</code> you have to specify them.
<code>genotFitness</code>	A matrix or data frame that contains explicitly the mapping of genotypes to fitness. For now, we only allow epistasis-like relations between genes (so you cannot code order effects this way). Genotypes can be specified in two ways: <ul style="list-style-type: none"> <li>• As a matrix (or data frame) with <math>g + 1</math> columns (where <math>g &gt; 1</math>). Each of the first <math>g</math> columns contains a 1 or a 0 indicating that the gene of that column is mutated or not. Column <math>g + 1</math> contains the fitness values. This is, for instance, the output you will get from <code>rfitness</code>. If the matrix has all columns named, those will be used for the names of the genes. Of course, except for column or row names, all entries in this matrix or data frame must be numeric.</li> <li>• As a two column data frame. The second column is fitness, and the first column are genotypes, given as a character vector. For instance, a row "A, B" would mean the genotype with both A and B mutated.</li> </ul> In all cases, fitness must be $\geq 0$ . If any possible genotype is missing, its fitness is assumed to be 1.
<code>keepInput</code>	If TRUE, whether to keep the original input. This is only useful for human consumption of the output. It is useful because it is easier to decode, say, the restriction table from the data frame than from the internal representation. But if you want, you can set it to FALSE and the object will be a little bit smaller.

## Details

`allFitnessEffects` is used for extremely flexible specification of fitness and mutator effects, including posets, XOR relationships, synthetic mortality and synthetic viability, arbitrary forms of epistasis, arbitrary forms of order effects, etc. Please, see the vignette for detailed and commented examples.

`allMutatorEffects` provide the same flexibility, but without order and posets (this might be included in the future, but I have seen no empirical or theoretical argument for their existence or relevance as of now, so I do not add them to minimize unneeded complexity).

If you use both for simulations in the same call to, say, `oncoSimulIndiv`, all the genes specified in `allMutatorEffects` MUST be included in the `allFitnessEffects` object. If you want to have genes that have no direct effect on fitness, but that affect mutation rate, you MUST specify them in the call to `allFitnessEffects`, for instance as `noIntGenes` with an effect of 0.

If you use `genotFitness` then you cannot pass modules, `noIntgenes`, epistasis, or `rT`. This makes sense, because using `genotFitness` is saying "this is the mapping of genotypes to fitness. Period", so we should not allow further modifications from other terms.

If you use `genotFitness` you need to be careful when you use Bozic's model (as you get a death rate of 0).

If you use `genotFitness` note that we force the WT (wildtype) to always be 1 so fitnesses are rescaled.

### Value

An object of class "fitnessEffects" or "mutatorEffects". This is just a list, but it is not intended for human consumption. The components are:

<code>long.rt</code>	The restriction table in "long format", so as to be easy to parse by the C++ code.
<code>long.epistasis</code>	Ditto, but for the epistasis specification.
<code>long.orderEffects</code>	Ditto for the order effects.
<code>long.geneNoInt</code>	Ditto for the non-interaction genes.
<code>geneModule</code>	Similar, for the gene-module correspondence.
<code>graph</code>	An <code>igraph</code> object that shows the restrictions, epistasis and order effects, and is useful for plotting.
<code>drv</code>	The numeric identifiers of the drivers. The numbers correspond to the internal numeric coding of the genes.
<code>rT</code>	If <code>keepInput</code> is TRUE, the original restriction table.
<code>epistasis</code>	If <code>keepInput</code> is TRUE, the original epistasis vector.
<code>orderEffects</code>	If <code>keepInput</code> is TRUE, the original order effects vector.
<code>noIntGenes</code>	If <code>keepInput</code> is TRUE, the original <code>noIntGenes</code> .

### Note

Please, note that the meaning of the fitness effects in the McFarland model is not the same as in the original paper; the fitness coefficients are transformed to allow for a simpler fitness function as a product of terms. This differs with respect to v.1. See the vignette for details.

The names of the genes and modules can be fairly arbitrary. But if you try hard you can confuse the parser. For instance, using gene or module names that contain ",", ":", or ">" is likely to get you into trouble. Of course, you know you should not try to use those characters because you know those characters have special meanings to separate names or indicate epistasis or order relationships. Right now, using those characters as names is caught (and result in stopping) if passed as names for `noIntGenes`.

### Author(s)

Ramon Diaz-Uriarte

### References

Diaz-Uriarte, R. (2015). Identifying restrictions in the order of accumulation of mutations during tumor progression: effects of passengers, evolutionary models, and sampling <http://www.biomedcentral.com/1471-2105/16/41/abstract>

McFarland, C.-D. et al. (2013). Impact of deleterious passenger mutations on cancer progression. *Proceedings of the National Academy of Sciences of the United States of America*, **110**(8), 2910–5.

**See Also**

[evalGenotype](#), [oncoSimulIndiv](#), [plot.fitnessEffects](#), [evalGenotypeFitAndMut](#), [rfitness](#), [plotFitnessLandscape](#)

**Examples**

```
## A simple poset or CBN-like example

cs <- data.frame(parent = c(rep("Root", 4), "a", "b", "d", "e", "c"),
                 child = c("a", "b", "d", "e", "c", "c", rep("g", 3)),
                 s = 0.1,
                 sh = -0.9,
                 typeDep = "MN")

cbn1 <- allFitnessEffects(cs)

plot(cbn1)

## A more complex example, that includes a restriction table
## order effects, epistasis, genes without interactions, and modules
p4 <- data.frame(parent = c(rep("Root", 4), "A", "B", "D", "E", "C", "F"),
                 child = c("A", "B", "D", "E", "C", "C", "F", "F", "G", "G"),
                 s = c(0.01, 0.02, 0.03, 0.04, 0.1, 0.1, 0.2, 0.2, 0.3, 0.3),
                 sh = c(rep(0, 4), c(-.9, -.9), c(-.95, -.95), c(-.99, -.99)),
                 typeDep = c(rep("--", 4),
                             "XMPN", "XMPN", "MN", "MN", "SM", "SM"))

oe <- c("C > F" = -0.1, "H > I" = 0.12)
sm <- c("I:J" = -1)
sv <- c("-K:M" = -.5, "K:-M" = -.5)
epist <- c(sm, sv)

modules <- c("Root" = "Root", "A" = "a1",
            "B" = "b1, b2", "C" = "c1",
            "D" = "d1, d2", "E" = "e1",
            "F" = "f1, f2", "G" = "g1",
            "H" = "h1, h2", "I" = "i1",
            "J" = "j1, j2", "K" = "k1, k2", "M" = "m1")

set.seed(1) ## for repeatability
noint <- rexp(5, 10)
names(noint) <- paste0("n", 1:5)

fea <- allFitnessEffects(rT = p4, epistasis = epist, orderEffects = oe,
                       noIntGenes = noint, geneToModule = modules)

plot(fea)

## Modules that show, between them,
## no epistasis (so multiplicative effects).
## We specify the individual terms, but no value for the ":".

fnme <- allFitnessEffects(epistasis = c("A" = 0.1,
                                       "B" = 0.2),
```

```

geneToModule = c("A" = "a1, a2",
                 "B" = "b1")

evalAllGenotypes(fnme, order = FALSE, addwt = TRUE)

## Epistasis for fitness and simple mutator effects

fe <- allFitnessEffects(epistasis = c("a : b" = 0.3,
                                     "b : c" = 0.5),
                      noIntGenes = c("e" = 0.1))

fm <- allMutatorEffects(noIntGenes = c("a" = 10,
                                       "c" = 5))

evalAllGenotypesFitAndMut(fe, fm, order = FALSE)

## Simple fitness effects (noIntGenes) and modules
## for mutators

fe2 <- allFitnessEffects(noIntGenes =
                        c(a1 = 0.1, a2 = 0.2,
                          b1 = 0.01, b2 = 0.3, b3 = 0.2,
                          c1 = 0.3, c2 = -0.2))

fm2 <- allMutatorEffects(epistasis = c("A" = 5,
                                       "B" = 10,
                                       "C" = 3),
                       geneToModule = c("A" = "a1, a2",
                                         "B" = "b1, b2, b3",
                                         "C" = "c1, c2"))

evalAllGenotypesFitAndMut(fe2, fm2, order = FALSE)

## Passing fitness directly, a complete fitness specification
## with a two column data frame with genotypes as character vectors

(m4 <- data.frame(G = c("A, B", "A", "WT", "B"), F = c(3, 2, 1, 4)))
fem4 <- allFitnessEffects(genotFitness = m4)

## Verify it interprets what it should: m4 is the same as the evaluation
## of the fitness effects (note row reordering)
evalAllGenotypes(fem4, addwt = TRUE, order = FALSE)

## Passing fitness directly, a complete fitness specification
## that uses a three column matrix

m5 <- cbind(c(0, 1, 0, 1), c(0, 0, 1, 1), c(1, 2, 3, 5.5))
fem5 <- allFitnessEffects(genotFitness = m5)

## Verify it interprets what it should: m5 is the same as the evaluation
## of the fitness effects
evalAllGenotypes(fem5, addwt = TRUE, order = FALSE)

```

```
## Passing fitness directly, an incomplete fitness specification
## that uses a three column matrix

m6 <- cbind(c(1, 1), c(1, 0), c(2, 3))
fem6 <- allFitnessEffects(genotFitness = m6)
evalAllGenotypes(fem6, addwt = TRUE, order = FALSE)

## Plotting a fitness landscape

fe2 <- allFitnessEffects(noIntGenes =
                        c(a1 = 0.1,
                          b1 = 0.01,
                          c1 = 0.3))

plot(evalAllGenotypes(fe2, order = FALSE))

## same as
plotFitnessLandscape(evalAllGenotypes(fe2, order = FALSE))

## same as
plotFitnessLandscape(fe2)

## Reinitialize the seed
set.seed(NULL)
```

---

benchmarks

*Summary results from some benchmarks reported in the vignette.*

---

## Description

Summary results from some benchmarks reported in the vignette. Included are timings, sizes of return objects and key output from each simulation.

They are here mainly to facilitate creation of table from the vignette itself. The scripts are available under "inst/miscell".

## Usage

```
data(benchmark_1)
data(benchmark_1_0.05)
data(benchmark_2)
data(benchmark_3)
```

## Format

Data frames.

**Examples**

```
data(benchmark_1)
benchmark_1
```

---

evalAllGenotypes	<i>Evaluate fitness/mutator effects of one or all possible genotypes.</i>
------------------	---

---

**Description**

Given a fitnessEffects/mutatorEffects description, obtain the fitness/mutator effects of a single or all genotypes.

**Usage**

```
evalGenotype(genotype, fitnessEffects, verbose = FALSE, echo = FALSE,
             model = "")

evalGenotypeMut(genotype, mutatorEffects, verbose = FALSE, echo = FALSE)

evalAllGenotypes(fitnessEffects, order = FALSE, max = 256, addwt = FALSE,
                 model = "")

evalAllGenotypesMut(mutatorEffects, max = 256, addwt = FALSE)

evalGenotypeFitAndMut(genotype, fitnessEffects,
                      mutatorEffects, verbose = FALSE, echo = FALSE,
                      model = "")

evalAllGenotypesFitAndMut(fitnessEffects, mutatorEffects,
                          order = FALSE, max = 256, addwt = FALSE,
                          model = "")
```

**Arguments**

genotype	(For evalGenotype). A genotype, as a character vector, with genes separated by "," or ">", or as a numeric vector. Use the same integers or characters used in the fitnessEffects object. This is a genotype in terms of genes, not modules. Using "," or ">" makes no difference: the sequence is always taken as the order in which mutations occurred. Whether order matters or not is encoded in the fitnessEffects object.
fitnessEffects	A fitnessEffects object, as produced by <a href="#">allFitnessEffects</a> .
mutatorEffects	A mutatorEffects object, as produced by <a href="#">allMutatorEffects</a> .
order	(For evalAllGenotypes). If TRUE, then order matters. If order matters, then generate not only all possible combinations of the genes, but all possible permutations for each combination.

max	(For evalAllGenotypes). By default, no output is shown if the number of possible genotypes exceeds the max. Increase as needed.
addwt	(For evalAllGenotypes). Add the wildtype (no mutations) explicitly?
model	Either nothing (the default) or "Bozic". If "Bozic" then the fitness effects contribute to decreasing the Death rate. Otherwise Birth rate is shown (and labeled as Fitness).
verbose	(For evalGenotype). If set to TRUE, print out the individual terms that are added to 1 (or subtracted from 1, if model is "Bozic").
echo	(For evalGenotype). If set to TRUE, show the input genotype and print out a message with the death rate or fitness value. Useful for some examples, as shown in the vignette.

### Value

For evalGenotype either the value of fitness or (if verbose = TRUE) the value of fitness and its individual components.

For evalAllGenotypes a data frame with two columns, the Genotype and the Fitness (or Death Rate, if Bozic). The notation for the Genotype column is as follows: when order does not matter, a comma "," separates the identifiers of mutated genes. When order matters, a genotype shown as "x > y \_ z" means that a mutation in "x" happened before a mutation in "y"; there is also a mutation in "z" (which could have happened before or after either of "x" or "y"), but "z" is a gene for which order does not matter. In all cases, a "WT" denotes the wild-type (or, actually, the genotype without any mutations).

If you use both fitnessEffects and mutatorEffects in a call, all the genes specified in mutatorEffects MUST be included in the fitnessEffects object. If you want to have genes that have no direct effect on fitness, but that affect mutation rate, you MUST specify them in the call to fitnessEffects, for instance as noIntGenes with an effect of 0.

### Note

Fitness is used in a slight abuse of the language. Right now, mutations contribute to the birth rate for all models except Bozic, where they modify the death rate. The general expression for fitness is the usual multiplicative one of  $\prod (1 + s_i)$ , where each  $s_i$  refers to the fitness effect of the given gene. When dealing with death rates, we use  $\prod (1 - s_i)$ .

Modules are, of course, taken into account if present (i.e., fitness is specified in terms of modules, but the genotype is specified in terms of genes).

About the naming. This is the convention used: "All" means we will go over all possible genotypes. A function that ends as "Genotypes" returns only fitness effects (for backwards compatibility and because mutator effects are not always used). A function that ends as "Genotype(s)Mut" returns only the mutator effects. A function that ends as "FitAndMut" will return both fitness and mutator effects.

Functions that return ONLY fitness or ONLY mutator effects are kept as separate functions because they free you from specifying mutator/fitness effects if you only want to play with one of them.

### Author(s)

Ramon Diaz-Uriarte

### See Also

[allFitnessEffects](#).

**Examples**

```

# A three-gene epistasis example
sa <- 0.1
sb <- 0.15
sc <- 0.2
sab <- 0.3
sbc <- -0.25
sabc <- 0.4

sac <- (1 + sa) * (1 + sc) - 1

E3A <- allFitnessEffects(epistasis =
  c("A:-B:-C" = sa,
    "-A:B:-C" = sb,
    "-A:-B:C" = sc,
    "A:B:-C" = sab,
    "-A:B:C" = sbc,
    "A:-B:C" = sac,
    "A : B : C" = sabc)
  )

evalAllGenotypes(E3A, order = FALSE, addwt = FALSE)
evalAllGenotypes(E3A, order = FALSE, addwt = TRUE, model = "Bozic")

evalGenotype("B, C", E3A, verbose = TRUE)

## Order effects and modules
ofe2 <- allFitnessEffects(orderEffects = c("F > D" = -0.3, "D > F" = 0.4),
  geneToModule =
  c("Root" = "Root",
    "F" = "f1, f2, f3",
    "D" = "d1, d2") )

evalAllGenotypes(ofe2, order = TRUE, max = 325)[1:15, ]

## Next two are identical
evalGenotype("d1 > d2 > f3", ofe2, verbose = TRUE)
evalGenotype("d1 , d2 , f3", ofe2, verbose = TRUE)

## This is different
evalGenotype("f3 , d1 , d2", ofe2, verbose = TRUE)
## but identical to this one
evalGenotype("f3 > d1 > d2", ofe2, verbose = TRUE)

## Restrictions in mutations as a graph. Modules present.

p4 <- data.frame(parent = c(rep("Root", 4), "A", "B", "D", "E", "C", "F"),
  child = c("A", "B", "D", "E", "C", "C", "F", "F", "G", "G"),
  s = c(0.01, 0.02, 0.03, 0.04, 0.1, 0.1, 0.2, 0.2, 0.3, 0.3),
  sh = c(rep(0, 4), c(-.9, -.9), c(-.95, -.95), c(-.99, -.99)),
  typeDep = c(rep("--", 4),
    "XMPN", "XMPN", "MN", "MN", "SM", "SM"))
fp4m <- allFitnessEffects(p4,
  geneToModule = c("Root" = "Root", "A" = "a1",
    "B" = "b1, b2", "C" = "c1",

```

```

"D" = "d1, d2", "E" = "e1",
"F" = "f1, f2", "G" = "g1"))

evalAllGenotypes(fp4m, order = FALSE, max = 1024, addwt = TRUE)[1:15, ]

evalGenotype("b1, b2, e1, f2, a1", fp4m, verbose = TRUE)

## Of course, this is identical; b1 and b2 are same module
## and order is not present here

evalGenotype("a1, b2, e1, f2", fp4m, verbose = TRUE)

evalGenotype("a1 > b2 > e1 > f2", fp4m, verbose = TRUE)

## We can use the exact same integer numeric id codes as in the
## fitnessEffects geneModule component:

evalGenotype(c(1L, 3L, 7L, 9L), fp4m, verbose = TRUE)

## Epistasis for fitness and simple mutator effects

fe <- allFitnessEffects(epistasis = c("a : b" = 0.3,
                                     "b : c" = 0.5),
                       noIntGenes = c("e" = 0.1))

fm <- allMutatorEffects(noIntGenes = c("a" = 10,
                                       "c" = 5))

evalAllGenotypesFitAndMut(fe, fm, order = "FALSE")

## Simple fitness effects (noIntGenes) and modules
## for mutators

fe2 <- allFitnessEffects(noIntGenes =
  c(a1 = 0.1, a2 = 0.2,
    b1 = 0.01, b2 = 0.3, b3 = 0.2,
    c1 = 0.3, c2 = -0.2))

fm2 <- allMutatorEffects(epistasis = c("A" = 5,
                                       "B" = 10,
                                       "C" = 3),
  geneToModule = c("A" = "a1, a2",
                  "B" = "b1, b2, b3",
                  "C" = "c1, c2"))

## Show only all the fitness effects
evalAllGenotypes(fe2, order = FALSE)

## Show only all mutator effects
evalAllGenotypesMut(fm2)

## Show all fitness and mutator
evalAllGenotypesFitAndMut(fe2, fm2, order = FALSE)

```

```
## This is probably not what you want
try(evalAllGenotypesMut(fe2))
## ... nor this
try(evalAllGenotypes(fm2))

## Show the fitness effect of a specific genotype
evalGenotype("a1, c2", fe2, verbose = TRUE)

## Show the mutator effect of a specific genotype
evalGenotypeMut("a1, c2", fm2, verbose = TRUE)

## Fitness and mutator of a specific genotype
evalGenotypeFitAndMut("a1, c2", fe2, fm2, verbose = TRUE)

## This is probably not what you want
try(evalGenotype("a1, c2", fm2, verbose = TRUE))

## Not what you want either
try(evalGenotypeMut("a1, c2", fe2, verbose = TRUE))
```

---

example-missing-drivers

*An example where there are intermediate missing drivers.*

---

## Description

An example where there are intermediate missing drivers. This is fictitious and I've never seen it. But it is here to check plots work even if there are no cases of some intermediate value of drivers (2 in this case). b11 contains the full, original data, whereas b12 contains the same data where there are no cases with exactly 2 drivers.

## Usage

```
data("ex_missing_drivers_b11"); data("ex_missing_drivers_b12")
```

## Format

Two objects of class "oncosimul".

## See Also

[plot.oncosimul](#)

## Examples

```
data(ex_missing_drivers_b11)
plot(ex_missing_drivers_b11, type = "line")
dev.new()
data(ex_missing_drivers_b12)
plot(ex_missing_drivers_b12, type = "line")
```



---

`examplesFitnessEffects`*Examples of fitness effects*

---

**Description**

Some examples fitnessEffects objects. This is a collection, in a list, of most of the fitnessEffects created (using [allFitnessEffects](#)) for the vignette. See the vignette for descriptions and references.

**Usage**

```
data("examplesFitnessEffects")
```

**Format**

The format is a list of fitnessEffects objects.

**See Also**

[allFitnessEffects](#)

**Examples**

```
data(examplesFitnessEffects)
plot(examplesFitnessEffects[["fea"]])
evalAllGenotypes(examplesFitnessEffects[["cbn1"]], order = FALSE)
```

---

`mcfLs`*mcfLs simulation from the vignette*

---

**Description**

Trimmed output from the simulation mcfLs in the vignette. This is a somewhat long run, and we have stored here the object (after trimming the Genotype matrix) to allow for plotting it.

**Usage**

```
data("mcfLs")
```

**Format**

An object of class "oncosimul2". A list.

**See Also**

[plot.oncosimul](#)

**Examples**

```
data(mcfLs)

plot(mcfLs, addtot = TRUE, lwdClone = 0.9, log = "")
summary(mcfLs)
```

---

oncoSimulIndiv	<i>Simulate tumor progression for one or more individuals, optionally returning just a sample in time.</i>
----------------	--

---

**Description**

Simulate tumor progression including possible restrictions in the order of driver mutations. Optionally add passenger mutations. Simulation is done using the BNB algorithm of Mather et al., 2012.

**Usage**

```
oncoSimulIndiv(fp, model = "Exp",
              numPassengers = 0, mu = 1e-6, muEF = NULL,
              detectionSize = 1e8, detectionDrivers = 4,
              detectionProb = NA,
              sampleEvery = ifelse(model %in% c("Bozic", "Exp"), 1,
                                   0.025),
              initSize = 500, s = 0.1, sh = -1,
              K = initSize/(exp(1) - 1), keepEvery = sampleEvery,
              minDetectDrvCloneSz = "auto",
              extraTime = 0,
              finalTime = 0.25 * 25 * 365, onlyCancer = TRUE,
              keepPhylog = FALSE,
              mutationPropGrowth = ifelse(model == "Bozic",
                                           FALSE, TRUE),
              max.memory = 2000, max.wall.time = 200,
              max.num.tries = 500,
              errorHitWallTime = TRUE,
              errorHitMaxTries = TRUE,
              verbosity = 0,
              initMutant = NULL,
              AND_DrvProbExit = FALSE,
              fixation = NULL,
              seed = NULL)

oncoSimulPop(Nindiv, fp, model = "Exp", numPassengers = 0, mu = 1e-6,
            muEF = NULL,
            detectionSize = 1e8, detectionDrivers = 4,
            detectionProb = NA,
            sampleEvery = ifelse(model %in% c("Bozic", "Exp"), 1,
                                  0.025),
            initSize = 500, s = 0.1, sh = -1,
            K = initSize/(exp(1) - 1), keepEvery = sampleEvery,
```

```

minDetectDrvCloneSz = "auto",
extraTime = 0,
finalTime = 0.25 * 25 * 365, onlyCancer = TRUE,
keepPhylog = FALSE,
mutationPropGrowth = ifelse(model == "Bozic",
                             FALSE, TRUE),
max.memory = 2000, max.wall.time = 200,
max.num.tries = 500,
errorHitWallTime = TRUE,
errorHitMaxTries = TRUE,
initMutant = NULL,
AND_DrvProbExit = FALSE,
fixation = NULL,
verbosity = 0,
mc.cores = detectCores(),
seed = "auto")

```

```

oncoSimulSample(Nindiv,
               fp,
               model = "Exp",
               numPassengers = 0,
               mu = 1e-6,
               muEF = NULL,
               detectionSize = round(runif(Nindiv, 1e5, 1e8)),

               detectionDrivers = {
                 if(inherits(fp, "fitnessEffects")) {
                   if(length(fp$drv)) {
                     nd <- (2: round(0.75 * length(fp$drv)))
                   } else {
                     nd <- 9e6
                   }
                 } else {
                   nd <- (2 : round(0.75 * max(fp)))
                 }
                 if (length(nd) == 1)
                   nd <- c(nd, nd)
                 sample(nd, Nindiv,
                       replace = TRUE)
               },
               detectionProb = NA,
               sampleEvery = ifelse(model %in% c("Bozic", "Exp"), 1,
                                     0.025),
               initSize = 500,
               s = 0.1,
               sh = -1,
               K = initSize/(exp(1) - 1),
               minDetectDrvCloneSz = "auto",
               extraTime = 0,
               finalTime = 0.25 * 25 * 365,
               onlyCancer = TRUE, keepPhylog = FALSE,

```

```

mutationPropGrowth = ifelse(model == "Bozic",
                             FALSE, TRUE),
max.memory = 2000,
max.wall.time.total = 600,
max.num.tries.total = 500 * Nindiv,
typeSample = "whole",
thresholdWhole = 0.5,
initMutant = NULL,
AND_DrvProbExit = FALSE,
fixation = NULL,
verbosity = 1,
showProgress = FALSE,
seed = "auto")

```

## Arguments

Nindiv	Number of individuals or number of different trajectories to simulate.
fp	Either a poset that specifies the order restrictions (see <a href="#">poset</a> if you want to use the specification as in v.1. Otherwise, a fitnessEffects object (see <a href="#">allFitnessEffects</a> ). Other arguments below (s, sh, numPassengers) make sense only if you use a poset, as they are included in the fitnessEffects object.
model	One of "Bozic", "Exp", "McFarlandLog" (the last one can be abbreviated to "McFL"). The default is "Exp".
numPassengers	This has no effect if you use the <a href="#">allFitnessEffects</a> specification. If you use the specification of v.1., the number of passenger genes. Note that using v.1 the total number of genes (drivers plus passengers) must be smaller than 64.  All driver genes should be included in the poset (even if they depend on no one and no one depends on them), and will be numbered from 1 to the total number of driver genes. Thus, passenger genes will be numbered from (number of driver genes + 1):(number of drivers + number of passengers).
mu	Mutation rate. Can be a single value or a named vector. If a single value, all genes will have the same mutation rate. If a named vector, the entries in the vector specify the gene-specific mutation rate. If you pass a vector, it must be named, and it must have entries for all the genes in the fitness specification. Passing a vector is only available when using fitnessEffects objects for fitness specification.  See also mutationPropGrowth.
muEF	Mutator effects. A mutatorEffects object as obtained from <a href="#">allMutatorEffects</a> . This specifies how mutations in certain genes change the mutation rate over all the genome. Therefore, this allows you to specify mutator phenotypes: models where mutation of one (or more) gene(s) leads to an increase in the mutation rate. This is only available for version 2 (and above) specifications.  All the genes specified in muEF MUST be included in fp. If you want to have genes that have no direct effect on fitness, but that affect mutation rate, you MUST specify them in fp, for instance as noIntGenes with an effect of 0.  If you use mutator effects you must also use fitnessEffects in fp.

detectionSize	<p>What is the minimal number of cells for cancer to be detected. For <code>oncoSimulSample</code> this can be a vector.</p> <p>If set to NA, <code>detectionSize</code> plays no role in stopping the simulations.</p>
detectionDrivers	<p>The minimal number of drivers (not modules, drivers, whether or not they are from the same module) present in any clone for cancer to be detected. For <code>oncoSimulSample</code> this can be a vector.</p> <p>For <code>oncoSimulSample</code>, if there are drivers (either because you are using a v.1 object or because you are using a <code>fitnessEffects</code> object with a <code>drvNames</code> component—see <a href="#">allFitnessEffects</a>—) the default is a vector of drivers from a uniform between 2 and 0.75 the total number of drivers. If there are no drivers (because you are using a <code>fitnessEffects</code> object without a <code>drvNames</code>, either because you specified it explicitly or because all of the genes are in the <code>noIntGenes</code> component) the simulations should not stop based on the number of drivers (and, thus, the default is set to 9e6).</p> <p>If set to NA, <code>detectionDrivers</code> plays no role in stopping the simulations.</p>
detectionProb	<p>Vector of arguments for the mechanism where probability of detection depends on size. If NA, this mechanism is not used. If ‘default’, the vector will be populated with default values. Otherwise, a named vector with some of the following named elements (see ‘Details’):</p> <ul style="list-style-type: none"> <li>• <code>PDBaseline</code>: Baseline size subtracted to total population size to compute the probability of detection. If not given explicitly, the default is <math>1.2 * \text{initSize}</math>.</li> <li>• <code>p2</code>: The probability of detection at population size <code>n2</code>. If you specify <code>p2</code> you must also specify <code>n2</code> and you must not specify <code>cPDetect</code>. The default is 0.1.</li> <li>• <code>n2</code>: The population size at which probability of detection is <code>p2</code>. The default is <math>2 * \text{initSize}</math>.</li> <li>• <code>cPDetect</code>: The change in probability of detection with size. If you specify it, you should not specify either of <code>p2</code> or <code>n2</code>. See ‘Details’.</li> <li>• <code>checkSizePEvery</code>: Time between successive checks for the probability of exiting as a function of population size. If not given explicitly, the default is 20. See ‘Details’.</li> </ul> <p>If you only provide some of the elements (except for the pair <code>p2</code>, <code>n2</code>, where you must provide both if you provide any), the rest will be filled with default values. This option can not be used with v.1 objects.</p>
sampleEvery	<p>How often the whole population is sampled. This is not the same as the interval between successive samples that are kept or stored (for that, see <code>keepEvery</code>).</p> <p>For very fast growing clones, you might need to have a small value here to minimize possible numerical problems (such as huge increase in population size between two successive samples that can then lead to problems for random number generators). Likewise, for models with density dependence (such as McF) this value should be very small.</p>
initSize	Initial population size.
K	Initial population equilibrium size in the McFarland models.
keepEvery	Time interval between successive whole population samples that are actually stored. This must be larger or equal to <code>sampleEvery</code> . If <code>keepEvery</code> is not a multiple integer of <code>sampleEvery</code> , the interval between successive samples that are stored will be the smallest multiple integer of <code>sampleEvery</code> that is larger than or equal to <code>keepEvery</code> .

If you want nice plots, set `sampleEvery` and `keepEvery` to small values (say, 5 or 2). Otherwise, you can use a `sampleEvery` of 1 but a `keepEvery` of 15, so that the return objects are not huge and the code runs a lot faster.

Setting `keepEvery = NA` means we only keep the very last sample. This is useful if you only care about the final state of the simulation, not its complete history.

#### `minDetectDrvCloneSz`

A value of 0 or larger than 0 (by default equal to `initSize` in the McFarland model). If larger than 0, when checking if we are done with a simulation, we verify that the sum of the population sizes of all clones that have a number of mutated drivers larger or equal to `detectionDrivers` is larger or equal to this `minDetectDrvCloneSz`.

The reason for this parameter is to ensure that, say, a clone with a certain number of drivers that would cause the simulation to end has not just appeared and is present in only one individual that might then immediately go extinct. This can be relevant in scenarios such as the McFarland model.

See also `extraTime`.

#### `extraTime`

A value larger than zero waits those many additional time periods before exiting after having reached the exit condition (population size, number of drivers).

The reason for this setting is to prevent the McFL models from always exiting at a time when one clone is increasing its size quickly (see `minDetectDrvCloneSz`). By setting an `extraTime` larger than 0, we can sample at points when we are at the plateau.

#### `finalTime`

What is the maximum number of time units that the simulation can run. Set to NA to disable this limit.

#### `onlyCancer`

Return only simulations that reach cancer?

If set to TRUE, only simulations that satisfy the `detectionDrivers` or the `detectionSize` requirements or that are "detected" because of the `detectionProb` mechanism will be returned: the simulation will be repeated, within the limits set by `max.num.tries` and `max.wall.time` (and, for `oncoSimulSample` also `max.num.tries.total` and `max.wall.time.total`), until one which meets the `detectionDrivers` or `detectionSize` or one which is detected stochastically under `detectionProb` is obtained.

If `onlyCancer = FALSE` the simulation is returned regardless of final population size or number of drivers in any clone and this includes simulations where the population goes extinct.

#### `keepPhylog`

If TRUE, keep track of when and from which clone each clone is created. See also [plotClonePhylog](#).

#### `mutationPropGrowth`

If TRUE, make mutation rate proportional to growth rate, so clones that grow faster also mutate faster. Thus,  $\text{mutation\_rate} = \mu * \text{birth\_rate}$ . This is a simple way of approximating that mutation happens at cell division (it is not strictly making mutation happen at cell division, since mutation is not strictly coupled with division). Of course, this only makes sense in models where birth rate changes.

#### `initMutant`

For v.2: a string with the mutations of the initial mutant, if any. This is the same format as for [evalGenotype](#). The default (if you pass nothing) is to start the simulation from the wildtype genotype with nothing mutated. For v.1 we no longer accept `initMutant`: it will be ignored.

<code>max.num.tries</code>	Only applies when <code>onlyCancer = TRUE</code> . What is the maximum number of times, for an individual simulation, we can repeat the simulation for it to reach cancer? There are certain parameter settings where reaching cancer is extremely unlikely and you might not want to run forever in those cases.
<code>max.num.tries.total</code>	Only applies when <code>onlyCancer = TRUE</code> and for <code>oncoSimulSample</code> . What is the maximum number of times, over all simulations for all individuals in a population sample, that we can repeat the simulations so that cancer is reached for all individuals? The idea is to set a limit on the average minimal probability of reaching cancer for a set of simulations to be accepted.
<code>max.wall.time</code>	Maximum wall time for the simulation of one individual (over all <code>max.num.tries</code> ). If the simulation is not done in this time, it is aborted.
<code>max.wall.time.total</code>	Maximum wall time for all the simulations (when using <code>oncoSimulSample</code> ), in seconds. If the simulation is not completed in this time, it is aborted. To prevent problems from a single individual simulation going wild, this limit is also enforced per simulation (so the run can be aborted directly from C++).
<code>errorHitMaxTries</code>	If <code>TRUE</code> (the default) a simulation that reaches the maximum number of repetitions allowed is considered not to have successfully finished and, thus, an error, and no output from it will be reported. This is often what you want. See <a href="#">Details</a> .
<code>errorHitWallTime</code>	If <code>TRUE</code> (the default) a simulation that reaches the maximum wall time is considered not to have successfully finished and, thus, an error, and no output from it will be reported. This is often what you want. See <a href="#">Details</a> .
<code>max.memory</code>	The largest size (in MB) of the matrix of Populations by Time. If it creating it would use more than this amount of memory, it is not created. This prevents you from accidentally passing parameters that will return an enormous object.
<code>verbosity</code>	If 0, run silently. Increasing values of verbosity provide progressively more information about intermediate steps, possible numerical notes/warnings from the C++ code, etc. Values less than 0 suppress some default notes: use with care.
<code>typeSample</code>	"singleCell" (or "single") for single cell sampling, where the probability of sampling a cell (a clone) is directly proportional to its population size. "wholeTumor" (or "whole") for whole tumor sampling (i.e., this is similar to a biopsy being the entire tumor). See <a href="#">samplePop</a> .
<code>thresholdWhole</code>	In whole tumor sampling, whether a gene is detected as mutated depends on <code>thresholdWhole</code> : a gene is considered mutated if it is altered in at least <code>thresholdWhole</code> proportion of the cells in that individual. See <a href="#">samplePop</a> .
<code>mc.cores</code>	Number of cores to use when simulating more than one individual (i.e., when calling <code>oncoSimulPop</code> ).
<code>showProgress</code>	If <code>TRUE</code> , provide information, during execution, of the individual done, and the number of attempts and time used.
<code>AND_DrvProbExit</code>	If <code>TRUE</code> , cancer will be considered to be reached if both the <code>detectionProb</code> mechanism and <code>detectionDrivers</code> are satisfied. This is an AND, not an OR condition. Using this option with <code>fixation</code> is not allowed (as it does not make much sense).

fixation	<p>If non-NULL, a list or a vector, where each element of is a string with a gene or a gene combination or a genotype (see below). Simulations will stop as soon as any of the genes or gene combinations or genotypes are fixed (i.e., reach a minimal frequency). If you pass gene combinations or genotypes, separate genes with commas (not '&gt;'); this means order is not (yet?) supported. This way of specifying gene combinations is the same as the one used for <code>initMutant</code> and <code>evalGenotype</code>.</p> <p>To differentiate between gene combinations and specific genotypes, genotypes are specified by prepending them with a "_". For instance, <code>fixation = c("A", "B, C")</code> specifies stopping on any genotypes with those gene combinations. In contrast, <code>fixation = c("_,A", "_,B, C" )</code> specifies stopping only on genotypes "A" or "B, C". See the vignette for further examples.</p> <p>In addition to the gene combinations or genotypes themselves, you can add to the list or vector the named elements <code>fixation_tolerance</code>, <code>min_successive_fixation</code> and <code>fixation_min_size</code>. <code>fixation_tolerance</code>: fixation is considered to have happened if the genotype/gene combinations specified as <code>genotypes/gene combinations</code> for fixation have reached a frequency <math>&gt; 1 - \text{fixation\_tolerance}</math>. (The default is 0, so we ask for genotypes/gene combinations with a frequency of 1, which might not be what you want with large mutation rates and complex fitness landscape with genotypes of similar fitness.). <code>min_successive_fixation</code>: during how many successive sampling periods the conditions of fixation need to be fulfilled before declaring fixation. These must be successive sampling periods without interruptions (i.e., a single period when the condition is not fulfilled will set the counter to 0). This can help to exclude short, transitional, local maxima that are quickly replaced by other genotypes. (The default is 50, but this is probably too small for "real life" usage). <code>fixation_min_size</code>: you might only want to consider fixation to have happened if a minimal size has been reached (this can help weed out local maxima that have fitness that is barely above that of the wild-type genotype). (The default is 0).</p> <p>Using this option with <code>AND_DrvProbExit</code> is not allowed (as it does not make much sense). This option is not allowed either with the old v.1 specification.</p>
s	<p>Selection coefficient for drivers. Only relevant if using a poset as this is included in the <code>fitnessEffects</code> object. This will eventually be deprecated.</p>
sh	<p>Selection coefficient for drivers with restrictions not satisfied. A value of 0 means there are no penalties for a driver appearing in a clone when its restrictions are not satisfied.</p> <p>To specify "sh=Inf" (in Diaz-Uriarte, 2015) use <code>sh = -1</code>.</p> <p>Only relevant if using a poset as this is included in the <code>fitnessEffects</code> object. This will eventually be deprecated.</p>
seed	<p>The seed for the C++ PRNG. You can pass a value. If you set it to NULL, then a seed will be generated in R and passed to C++. If you set it to "auto", then if you are using v.1, the behavior is the same as if you set it to NULL (a seed will be generated in R and passed to C++) but if you are using v.2, a random seed will be produced in C++. If you need reproducibility, either pass a value or set it to NULL (setting it to NULL will make the C++ seed reproducible if you use the same seed in R via <code>set.seed</code>). However, even using the same value of seed is unlikely to give the exact same results between platforms and compilers. Moreover, note that the defaults for seed are not the same in <code>oncoSimulIndiv</code>, <code>oncoSimulPop</code> and <code>oncoSimulSample</code>.</p> <p>When using <code>oncoSimulPop</code>, if you want reproducibility, you might want to, in addition to setting <code>seed = NULL</code>, also do <code>RNGkind("L'Ecuyer-CMRG")</code> as we</p>

use `mclapply`; look at the vignette of `parallel`.

## Details

The basic simulation algorithm implemented is the BNB one of Mather et al., 2012, where I have added modifications to fitness based on the restrictions in the order of mutations.

Full details about the algorithm are provided in Mather et al., 2012. The evolutionary models, including references, and the rest of the parameters are explained in Diaz-Uriarte, 2014, especially in the Supplementary Material. The model called "Bozic" is based on Bozic et al., 2010, and the model called "McFarland" in McFarland et al., 2013.

`oncoSimulPop` simply calls `oncoSimulIndiv` multiple times. When run on POSIX systems, it can use multiple cores (via `mclapply`).

The summary methods for these classes return some of the return values (see next) as a one-row (for class `oncosimul`) or multiple row (for class `oncosimulpop`) data frame. The `print` methods for these classes simply print the summary.

Changing options `errorHitMaxTries` and `errorHitWallTime` can be useful when conducting many simulations, as in the call to `oncoSimulPop`: setting them to `TRUE` means nothing is recorded for those simulations where ending conditions are not reached but setting them to `FALSE` would allow you to record the output; this would potentially result in a mixture where some simulations would not have reached the ending condition, but this might sometimes be what you want. Note, however, that `oncoSimulSample` always has both them to `TRUE`, as it could not be otherwise.

`GenotypesWDistinctOrderEff` provides the information about order effects that is missing from `Genotypes`. When there are order effects, the `Genotypes` matrix can contain genotypes that are not distinguishable. Suppose there are two genes, the first and the second. In the `Genotype` output you can get two columns where there is a 1 in both genes: those two columns correspond to the two possible orders (first gene mutated first, or first gene mutated after the second). `GenotypesWDistinctOrderEff` disambiguates this. The same is done by `GenotypesLabels`; this is easier to decode for a human (a string of gene labels) but a little bit harder to parse automatically. Note that when you use the default `print` method for this object, you get, among others, a two-column display with the `GenotypeLabels` information. When order matters, a genotype shown as "x > y \_ z" means that a mutation in "x" happened before a mutation in "y"; there is also a mutation in "z" (which could have happened before or after either of "x" or "y"), but "z" is a gene for which order does not matter. When order does not matter, a comma "," separates the identifiers of mutated genes.

Detection of cancer can be a deterministic process, where cancer is always detected (and, thus, simulation ended) when certain conditions are met (`detectionSize`, `detectionDrivers`, `fixation`). Alternatively, it can be stochastic process where probability of detection depends on size. Every so often (see below) we assess population size, and detect cancer or not probabilistically (comparing the probability of detection for that size with a random uniform number). Probability of detection changes with population size according to the function

$$1 - e^{-cPDetect((populationsize - PDBaseline)/PDBaseline)}$$

You can pass `cPDetect` manually (you will need to set `n2` and `p2` to `NA`). However, it might be more intuitive to specify the pair `n2`, `p2`, such that the probability of detection is `p2` for population size `n2` (and from that pair we solve for the value of `cPDetect`). How often do we check? That is controlled by `checkSizePEvery`, the (minimal) time between successive checks (from among the sampling times given by `sampleEvery`: the interval between successive assessments will be the smallest multiple integer of `sampleEvery` that is larger than `checkSizePEvery` —see vignette for details).

checkSizePEvery has, by default, a different (and much larger) value than sampleEvery both to allow to examine the effects of sampling, and to avoid many costly random number generations.

Please note that detectionProb is NOT available with version 1 objects.

## Value

For oncoSimulIndiv a list, of class "oncosimul", with the following components:

pops.by.time	A matrix of the population sizes of the clones, with clones in columns and time in row. Not all clones are shown here, only those that were present in at least one of the keepEvery samples.
NumClones	Total number of clones in the above matrix. This is not the total number of distinct clones that have appeared over all simulations (which is likely to be larger or much larger).
TotalPopSize	Total population size at the end.
Genotypes	A matrix of genotypes. For each of the clones in the pops.by.time matrix, its genotype, with a 0 if the gene is not mutated and a 1 if it is mutated.
MaxNumDrivers	The largest number of mutated driver genes ever seen in the simulation in any clone.
MaxDriversLast	The largest number of mutated drivers in any clone at the end of the simulation.
NumDriversLargestPop	The number of mutated driver genes in the clone with largest population size.
LargestClone	Population size of the clone with largest number of population size.
PropLargestPopLast	Ratio of LargestClone/TotalPopSize
FinalTime	The time (in time units) at the end of the simulation.
NumIter	The number of iterations of the BNB algorithm.
HittedWallTime	TRUE if we reached the limit of max.wall.time. FALSE otherwise.
TotalPresentDrivers	The total number of mutated driver genes, whether or not in the same clone. The number of elements in OccurringDrivers, below.
CountByDriver	A vector of length number of drivers, with the count of the number of clones that have that driver mutated.
OccurringDrivers	The actual number of drivers mutated.
PerSampleStats	A 5 column matrix with a row for each sampling period. The columns are: total population size, population size of the largest clone, the ratio of the two, the largest number of drivers in any clone, and the number of drivers in the clone with the largest population size.
other	A list that contains statistics for an estimate of the simulation error when using the McFarland model as well as other statistics. For the McFarland model, the relevant value is errorMF, which is -99 unless in the McFarland model. For the McFarland model it is the largest difference of successive death rates. The entries named minDMratio and minBMratio are the smallest ratio, over all simulations, of death rate to mutation rate and birth rate to mutation rate, respectively. The BNB algorithm thrives when those are large.

For oncoSimulPop a list of length Nindiv, and of class "oncosimulpop", where each element of the list is itself a list, of class oncosimul, with components as described above.

In v.2, the output is of both class "oncosimul" and "oncosimul2". The oncoSimulIndiv return object differs in

#### GenotypesWDistinctOrderEff

A list of vectors, where each vector corresponds to a genotype in the Genotypes, showing (where it matters) the order of mutations. Each vector shows the genotypes, with the numeric codes, showing explicitly the order when it matters. So if you have genes 1, 2, 7 for which order relationships are given, and genes 3, 4, 5, 6 for which other interactions exist, any mutations in 1, 2, 7 are shown first, and in the order they occurred, before showing the rest of the mutations. See details.

#### GenotypesLabels

The genotypes, as character vectors with the original labels provided (i.e., not the integer codes). As before, mutated genes, for those where order matters, come first, and are separated by the rest by a "\_". See details.

#### OccurringDrivers

This is the same as in v.1, but we use the labels, not the numeric id codes. Of course, if you entered integers as labels for the genes, you will see numbers (however, as a character string).

### Note

Please, note that the meaning of the fitness effects in the McFarland model is not the same as in the original paper; the fitness coefficients are transformed to allow for a simpler fitness function as a product of terms. This differs with respect to v.1. See the vignette for details.

### Author(s)

Ramon Diaz-Uriarte

### References

Bozic, I., et al., (2010). Accumulation of driver and passenger mutations during tumor progression. *Proceedings of the National Academy of Sciences of the United States of America*, **107**, 18545–18550.

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Gerstung et al., 2011. The Temporal Order of Genetic and Pathway Alterations in Tumorigenesis. *PLoS ONE*, 6.

McFarland, C.-D. et al. (2013). Impact of deleterious passenger mutations on cancer progression. *Proceedings of the National Academy of Sciences of the United States of America*, **110**(8), 2910–5.

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### See Also

[plot.oncosimul](#), [examplePosets](#), [samplePop](#), [allFitnessEffects](#)

**Examples**

```
#####
#####
##### Examples using v.1
#####
#####

## use poset p701
data(examplePosets)
p701 <- examplePosets[["p701"]]

## Exp Model

b1 <- oncoSimulIndiv(p701)
summary(b1)

plot(b1, addtot = TRUE)

## McFarland; use a small sampleEvery, but also a reasonable
## keepEvery.
## We also modify mutation rate to values similar to those in the
## original paper.
## Note that detectionSize will play no role
## finalTime is large, since this is a slower process
## initSize is set to 4000 so the default K is larger and we are likely
## to reach cancer. Alternatively, set K = 2000.

m1 <- oncoSimulIndiv(p701,
                    model = "McFL",
                    mu = 5e-7,
                    initSize = 4000,
                    sampleEvery = 0.025,
                    finalTime = 15000,
                    keepEvery = 10,
                    onlyCancer = FALSE)
plot(m1, addtot = TRUE, log = "")

## Simulating 4 individual trajectories
## (I set mc.cores = 2 to comply with --as-cran checks, but you
## should either use a reasonable number for your hardware or
## leave it at its default value).

p1 <- oncoSimulPop(4, p701,
                  keepEvery = 10,
                  mc.cores = 2)

summary(p1)
samplePop(p1)

p2 <- oncoSimulSample(4, p701)
```

```
#####
#####
##### Examples using v.2:
#####
#####

#### A model similar to the one in McFarland. We use 2070 genes.

set.seed(456)
nd <- 70
np <- 2000
s <- 0.1
sp <- 1e-3
spp <- -sp/(1 + sp)
mcf1 <- allFitnessEffects(noIntGenes = c(rep(s, nd), rep(spp, np)),
                          drv = seq.int(nd))
mcf1s <- oncoSimulIndiv(mcf1,
                       model = "McFL",
                       mu = 1e-7,
                       detectionSize = 1e8,
                       detectionDrivers = 100,
                       sampleEvery = 0.02,
                       keepEvery = 2,
                       initSize = 2000,
                       finalTime = 1000,
                       onlyCancer = FALSE)
plot(mcf1s, addtot = TRUE, lwdClone = 0.6, log = "")
summary(mcf1s)
plot(mcf1s)

#### Order effects with modules, and 5 genes without interactions
#### with fitness effects from an exponential distribution

oi <- allFitnessEffects(orderEffects =
                        c("F > D" = -0.3, "D > F" = 0.4),
                        noIntGenes = rexp(5, 10),
                        geneToModule =
                          c("Root" = "Root",
                            "F" = "f1, f2, f3",
                            "D" = "d1, d2") )
oiI1 <- oncoSimulIndiv(oi, model = "Exp")
oiI1$GenotypesLabels
oiI1 ## note the order and separation by "_"

oiP1 <- oncoSimulPop(2, oi,
                    keepEvery = 10,
                    mc.cores = 2)

summary(oiP1)

## Even if order exists, this cannot reflect it;
## G1 to G10 are d1, d2, f1..,f3, and the 5 genes without
```

```

## interaction
samplePop(oiP1)

oiS1 <- oncoSimulSample(2, oi)

## The output contains only the summary of the runs AND
## the sample:
oiS1

## And their sizes do differ
object.size(oiS1)
object.size(oiP1)

##### Using a poset for pancreatic cancer from Gerstung et al.
###      (s and sh are made up for the example; only the structure
###      and names come from Gerstung et al.)

pancr <- allFitnessEffects(data.frame(parent = c("Root", rep("KRAS", 4), "SMAD4", "CDNK2A",
      "TP53", "TP53", "MLL3"),
      child = c("KRAS", "SMAD4", "CDNK2A",
      "TP53", "MLL3",
      rep("PXDN", 3), rep("TGFB2", 2)),
      s = 0.05,
      sh = -0.3,
      typeDep = "MN"))

plot(pancr)

### Use an exponential growth model

pancr1 <- oncoSimulIndiv(pancr, model = "Exp")
pancr1
summary(pancr1)
plot(pancr1)
pancr1$GenotypesLabels

## Pop and Sample
pancrPop <- oncoSimulPop(4, pancr,
      keepEvery = 10,
      mc.cores = 2)
summary(pancrPop)
pancrSPop <- samplePop(pancrPop)
pancrSPop

pancrSamp <- oncoSimulSample(2, pancr)
pancrSamp

## Using gene-specific mutation rates
muv <- c("U" = 1e-3, "z" = 1e-7, "e" = 1e-6, "m" = 1e-5, "D" = 1e-4)
ni <- rep(0.01, 5)
names(ni) <- names(muv)

```

```
femuv <- allFitnessEffects(noIntGenes = ni)
oncoSimulIndiv(femuv, mu = muv)
```

```
## Reinitialize the RNG
set.seed(NULL)
```

---

OncoSimulWide2Long	<i>Convert the pops.by.time component of an oncosimul object into "long" format.</i>
--------------------	--

---

### Description

Conver the pops.by.time component from its "wide" format (with one column for time, and as many columns as clones/genotypes) into "long" format, so that it can be used with other functions, for instance for plots.

### Usage

```
OncoSimulWide2Long(x)
```

### Arguments

x                    An object of class oncosimul or oncosimul2.

### Value

A data frame with four columns: Time; Y, the number of cells (the population size); Drivers, a factor with the number of drivers of the given genotype; Genotype, the genotyp.

### Author(s)

Ramon Diaz-Uriarte

### See Also

[oncoSimulIndiv](#)

### Examples

```
data(examplePosets)
## An object of class oncosimul
p705 <- examplePosets[["p705"]]
p1 <- oncoSimulIndiv(p705)
class(p1)
lp1 <- OncoSimulWide2Long(p1)
head(lp1)
summary(lp1)

## An object of class oncosimul2
data(examplesFitnessEffects)

sm <- oncoSimulIndiv(examplesFitnessEffects$cbn1,
```

```

      model = "McFL",
      mu = 5e-7,
      detectionSize = 1e8,
      detectionDrivers = 2,
      sampleEvery = 0.025,
      keepEvery = 5,
      initSize = 2000,
      onlyCancer = FALSE)

class(sm)
lsm <- OncoSimulWide2Long(sm)
head(lsm)
summary(lsm)

```

---

plot.fitnessEffects    *Plot fitnessEffects objects.*

---

### Description

Plot the restriction table/graph of restrictions, the epistasis, and the order effects in a fitnessEffects object. This is not a plot of the fitness landscape; for that, see [plotFitnessLandscape](#).

### Usage

```

## S3 method for class 'fitnessEffects'
plot(x, type = "graphNEL", layout = NULL,
     expandModules = FALSE, autofit = FALSE,
     scale_char = ifelse(type == "graphNEL", 1/10, 5),
     return_g = FALSE, lwdf = 1, ...)

```

### Arguments

x	A fitnessEffects object, as produced by <a href="#">allFitnessEffects</a> .
type	Whether you want a "graphNEL" or an "igraph" graph.
layout	For "igraph", the layout. For example, if you know you really have only a tree you might want to use <code>layout.reingold.tilford</code> . Note that there is very limited support for passing options, etc. In most cases, it is either the default or the <code>layout.reingold.tilford</code> .
expandModules	If there are modules with multiple genes, if you set this to TRUE modules will be replaced by their genes.
autofit	If TRUE, we try to fit the edges to the labels. This is a very experimental feature, likely to be not very robust.
scale_char	If using <code>autofit = TRUE</code> , the scaling factor for the size of the rectangles as a function of the number of characters. You have to play with this because the best value can depend on a number of things.
return_g	If TRUE, the graph object (graphNEL or igrap) is returned.
lwdf	The multiplier factor for lwd when using "graphNEL".
...	Other arguments passed to plot. Not used for now.

**Value**

A plot.

Order and epistatic relationships have orange edges. OR (semimonotone) relationships blue, and XOR red. All others have black edges (so AND and unique edges from root). Epistatic relationships, being symmetrical, have no arrows between nodes and have a dotted line type. Order relationships have an arrow from the earlier to the later event and have a different dotted line (lty 3).

If return\_g is TRUE, you are returned also the graph object (igraph or graphNEL) so that you can manipulate it further.

**Note**

The purpose of the plot is to get a quick idea of the relationships. Note that three-way (or higher order) epistatic relationships cannot be shown as such (we would show all possible pairs, but that is not quite the same thing). Likewise, there is no reasonable way to convey the presence of a "-" in the epistatic relationship.

Genes without interactions are not shown.

**Author(s)**

Ramon Diaz-Uriarte

**See Also**

[allFitnessEffects](#), [plotFitnessLandscape](#)

**Examples**

```
cs <- data.frame(parent = c(rep("Root", 4), "a", "b", "d", "e", "c"),
  child = c("a", "b", "d", "e", "c", "c", rep("g", 3)),
  s = 0.1,
  sh = -0.9,
  typeDep = "MN")

cbn1 <- allFitnessEffects(cs)
plot(cbn1, "igraph")

library(igraph) ## to make layouts available
plot(cbn1, "igraph", layout = layout.reingold.tilford)

### A DAG with the three types of relationships
p3 <- data.frame(parent = c(rep("Root", 4), "a", "b", "d", "e", "c", "f"),
  child = c("a", "b", "d", "e", "c", "c", "f", "f", "g", "g"),
  s = c(0.01, 0.02, 0.03, 0.04, 0.1, 0.1, 0.2, 0.2, 0.3, 0.3),
  sh = c(rep(0, 4), c(-.9, -.9), c(-.95, -.95), c(-.99, -.99)),
  typeDep = c(rep("--", 4),
    "XMPN", "XMPN", "MN", "MN", "SM", "SM"))
fp3 <- allFitnessEffects(p3)

plot(fp3)

plot(fp3, "igraph", layout = layout.reingold.tilford)
```

```

## A more complex example, that includes a restriction table
## order effects, epistasis, genes without interactions, and modules
p4 <- data.frame(parent = c(rep("Root", 4), "A", "B", "D", "E", "C", "F"),
  child = c("A", "B", "D", "E", "C", "C", "F", "F", "G", "G"),
  s = c(0.01, 0.02, 0.03, 0.04, 0.1, 0.1, 0.2, 0.2, 0.3, 0.3),
  sh = c(rep(0, 4), c(-.9, -.9), c(-.95, -.95), c(-.99, -.99)),
  typeDep = c(rep("--", 4),
    "XMPN", "XMPN", "MN", "MN", "SM", "SM"))

oe <- c("C > F" = -0.1, "H > I" = 0.12)
sm <- c("I:J" = -1)
sv <- c("-K:M" = -.5, "K:-M" = -.5)
epist <- c(sm, sv)

modules <- c("Root" = "Root", "A" = "a1",
  "B" = "b1, b2", "C" = "c1",
  "D" = "d1, d2", "E" = "e1",
  "F" = "f1, f2", "G" = "g1",
  "H" = "h1, h2", "I" = "i1",
  "J" = "j1, j2", "K" = "k1, k2", "M" = "m1")

noint <- rexp(5, 10)
names(noint) <- paste0("n", 1:5)

fea <- allFitnessEffects(rT = p4, epistasis = epist, orderEffects = oe,
  noIntGenes = noint, geneToModule = modules)

plot(fea)
plot(fea, expandModules = TRUE)
plot(fea, type = "igraph")

```

---

plot.oncosimul

*Plot simulated tumor progression data.*

---

## Description

Plots data generated from the simulations, either for a single individual or for a population of individuals, with time units in the x axis and number of cells in the y axis.

In "drivers" plots, by default, all clones with the same number of drivers are plotted using the same colour (but different line types), and clones with different number of drivers are plotted in different colours. Plots can alternatively display genotypes instead of drivers.

Plots available are line plots, stacked area, and stream plots.

## Usage

```

## S3 method for class 'oncosimul'
plot(x,
      show = "drivers",
      type = ifelse(show == "genotypes",

```

```

                                "stacked", "line"),
col = "auto",
log = ifelse(type == "line", "y", ""),
ltyClone = 2:6,
lwdClone = 0.9,
ltyDrivers = 1,
lwdDrivers = 3,
xlab = "Time units",
ylab = "Number of cells",
plotClones = TRUE,
plotDrivers = TRUE,
addtot = FALSE,
addtotlwd = 0.5,
ylim = NULL,
xlim = NULL,
thinData = FALSE,
thinData.keep = 0.1,
thinData.min = 2,
plotDiversity = FALSE,
order.method = "as.is",
stream.center = TRUE,
stream.frac.rand = 0.01,
stream.spar = 0.2,
border = NULL,
lwdStackedStream = 1,
srange = c(0.4, 1),
vrage = c(0.8, 1),
breakSortColors = "oe",
legend.ncols = "auto", ...)

## S3 method for class 'oncosimulpop'
plot(x,

      ask = TRUE,
      show = "drivers",
      type = ifelse(show == "genotypes",
                    "stacked", "line"),
      col = "auto",
      log = ifelse(type == "line", "y", ""),
      ltyClone = 2:6,
      lwdClone = 0.9,
      ltyDrivers = 1,
      lwdDrivers = 3,
      xlab = "Time units",
      ylab = "Number of cells",
      plotClones = TRUE,
      plotDrivers = TRUE,
      addtot = FALSE,
      addtotlwd = 0.5,
      ylim = NULL,
      xlim = NULL,
      thinData = FALSE,
      thinData.keep = 0.1,

```

```

thinData.min = 2,
plotDiversity = FALSE,
order.method = "as.is",
stream.center = TRUE,
stream.frac.rand = 0.01,
stream.spar = 0.2,
border = NULL,
lwdStackedStream = 1,
srange = c(0.4, 1),
vrangle = c(0.8, 1),
breakSortColors = "oe",
legend.ncols = "auto",
...)
```

### Arguments

x	An object of class <code>oncosimul</code> (for <code>plot.oncosimul</code> ) or <code>oncosimulpop</code> (for <code>plot.oncosimulpop</code> ).
ask	Same meaning as in <code>par</code> .
show	One of "drivers" or "genotypes". If "drivers" the legend will reflect the number of drivers. If "genotypes" you will be shown genotypes. You probably want to limit "genotypes" to those cases where only a relatively small number of genotypes exist (or the plot will be an unmanageable mess). The default is "drivers".
type	One of "line", "stacked", "stream". If "line", you are shown lines for each genotype or clone. This means that to get an idea of the total population size you need to use <code>plotDrivers = TRUE</code> with <code>addtot = TRUE</code> , or do the visual calculation in your head. If "stacked" a stacked area plot. If "stream" a stream plot. Since these stack areas, you immediately get the total population. But that also means you cannot use <code>log</code> . The default is to use "line" for <code>show = "drivers"</code> and "stacked" for <code>show = "genotypes"</code> .
col	Colour of the lines/areas. For <code>show = "drivers"</code> each type of clone (where type is defined by number of drivers) has a different color. For <code>show = "genotypes"</code> color refers to genotypes. The vector is recycled as needed. The default is "auto". If you have <code>show == "genotypes"</code> we start from the "Dark2" palette from <code>brewer.pal</code> in the <code>RColorBrewer</code> package and extend the palette via <code>colorRampPalette</code> . For <code>show == "drivers"</code> and <code>type == "line"</code> we use a vector of eight colors (that are, then recycled as needed). If you use "stacked" or "stream", however, instead of "line", then we generate colors via a HSV specification that tries to: a) make it easy to differentiate between different drivers (by not having like colors for adjacent numbers of drivers); b) make it easy to have a "representative" driver color while using slightly different colors for different clones of a driver. See the code by doing <code>OncoSimulR:::myhsvcols</code> . You can specify your own vector of colors, but it will be ignored with <code>show == "drivers"</code> .
log	See log in <code>plot.default</code> . The default is to have "y" for <code>type == "line"</code> , and that will make the y axis logarithmic. Stacked and stream area plots do not allow for logarithmic y axis (since those depend on the additivity of areas but $\log(a + b) \neq \log(a) + \log(b)$ ).

ltyClone	Line type for each clone. Recycled as needed. You probably do not want to use lty=1 for any clone, to differentiate from the clone type, unless you change the setting for ltyDrivers.
lwdClone	Line width for clones.
ltyDrivers	Line type for the driver type.
lwdDrivers	Line width for the driver type.
xlab	Same as xlab in <a href="#">plot.default</a> .
ylab	Same as ylab in <a href="#">plot.default</a> .
plotClones	Should clones be plotted?
plotDrivers	Should clone types (which are defined by number of drivers), be plotted? (Only applies when using show = "drivers").
addtot	If TRUE, add a line with the total population size.
addtotlwd	Line width for total population size.
ylim	If non NULL, limits of the y axis. Same as in <a href="#">plot.default</a> . If NULL, the limits are calculated automatically.
xlim	If non NULL, limits of the x axis. Same as in <a href="#">plot.default</a> . If NULL, the limits are calculated automatically. Using a non-NULL range smaller than the range of observed values of time can also lead to speed ups of large figures (since we trim the data).
thinData	<p>If TRUE, the data plotted is a subset of the original data. The original data are "thinned" in such a way that the origin of each clone is not among the non-shown data (i.e., so that we can see when each clone/driver originates).</p> <p>Thinning is done to reduce the plot size and to speed up plotting.</p> <p>Note that thinning is carried out before dealing with the plot axis, so the actual number of points to be plotted could be a lot less (if you reduce the x-axis considerably) than those returned from the thinning. (In extreme cases this could lead to crashes when trying to use stream plots if, say, you end up plotting only three values).</p>
thinData.keep	The fraction of the data to keep (actually, a lower bound on the fraction of data to keep).
thinData.min	Any time point for which a clone has a population size > thinData.min will be kept (i.e., will not be removed from) in the data.
plotDiversity	<p>If TRUE, we also show, on top of the main figure, Shannon's diversity index (and we consider as distinct those genotypes with different order of mutations when order matters).</p> <p>If you set this to true, using <code>par(mfrow = c(2, 2))</code> and similar will not work (since we use <code>par(fig = )</code> to display the diversity as the top plot).</p>
order.method	For stacked and stream plots. <code>c("as.is", "max", "first")</code> . "as.is": plot in order of y column; "max": plot in order of when each y series reaches maximum value. "first": plot in order of when each y series first value > 0.
stream.center	For stream plots. If TRUE, the stacked polygons will be centered so that the middle, i.e. baseline ("g0"), of the stream is approximately equal to zero. Centering is done before the addition of random wiggle to the baseline.
stream.frac.rand	For stream plots. Fraction of the overall data "stream" range used to define the range of random wiggle (uniform distribution) to be added to the baseline 'g0'.

stream.spar	Setting for smooth.spline function to make a smoothed version of baseline "g0".
border	For stacked and stream plots. Border colors for polygons corresponding to y columns (will recycle) (see <a href="#">polygon</a> for details).
lwdStackedStream	border line width for polygons corresponding to y columns (will recycle).
srange	Range of values of s in the HSV specification of colors (see col for details. Only applies when using "stacked" or "stream" plots and col == "auto".)
vrange	Range of values of v in the HSV specification of colors (see col for details. Only applies when using "stacked" or "stream" plots and col == "auto".)
breakSortColors	How to try to minimize that similar colors be used for contiguous or nearby driver categories. The default is "oe" which resorts them in alternating way. The other two options are "distave", where we alternate after folding from the mean and "random" where the colors are randomly sorted. Only applies when using "stacked" or "stream" plots and col == "auto".
legend.ncols	The number of columns of the legend. If "auto" (the default), will have one column for six or less entries, and two for more than six.
...	Other arguments passed to plots. For instance, main.

**Author(s)**

Ramon Diaz-Uriarte. Marc Taylor for stacked and stream plots.

**See Also**

[oncoSimulIndiv](#)

**Examples**

```
data(examplePosets)
p701 <- examplePosets[["p701"]]

## Simulate and plot a single individual, including showing
## Shannon's diversity index
b1 <- oncoSimulIndiv(p701)
plot(b1, addtot = TRUE, plotDiversity = TRUE)

## A stacked area plot
plot(b1, type = "stacked", plotDiversity = TRUE)

## And what if I show a stream plot?
plot(b1, type = "stream", plotDiversity = TRUE)

## Simulate and plot 2 individuals
## (I set mc.cores = 2 to comply with --as-cran checks, but you
## should either use a reasonable number for your hardware or
## leave it at its default value).

p1 <- oncoSimulPop(2, p701, mc.cores = 2)

par(mfrow = c(1, 2))
plot(p1, ask = FALSE)
```

```

## Stacked; we cannot log here, and harder to see patterns
plot(p1, ask = FALSE, type = "stacked")

## Show individual genotypes and drivers for an
## epistasis case with at most eight genotypes

sa <- 0.1
sb <- -0.2
sab <- 0.25
sac <- -0.1
sbc <- 0.25
sv2 <- allFitnessEffects(epistasis = c("-A : B" = sb,
                                       "A : -B" = sa,
                                       "A : C" = sac,
                                       "A:B" = sab,
                                       "-A:B:C" = sbc),
                        geneToModule = c(
                          "Root" = "Root",
                          "A" = "a1, a2",
                          "B" = "b",
                          "C" = "c"))
evalAllGenotypes(sv2, order = FALSE, addwt = TRUE)
e1 <- oncoSimulIndiv(sv2, model = "McFL",
                    mu = 5e-6,
                    sampleEvery = 0.02,
                    keepEvery = 1,
                    initSize = 2000,
                    finalTime = 3000,
                    onlyCancer = FALSE)

## Drivers and clones
plot(e1, show = "drivers")

## Make genotypes explicit
plot(e1, show = "genotypes")

## Oh, but I want other colors
plot(e1, show = "genotypes", col = rainbow(8))

## and actually I want a line plot
plot(e1, show = "genotypes", type = "line")

```

---

plotClonePhylog

*Plot a parent-child relationship of the clones.*


---

### Description

Plot a parent-child relationship of the clones, controlling which clones are displayed, and whether to shown number of times of appearance, and time of first appearance of a clone.

**Usage**

```
plotClonePhylog(x, N = 1, t = "last", timeEvents = FALSE,
               keepEvents = FALSE, fixOverlap = TRUE,
               returnGraph = FALSE, ...)
```

**Arguments**

x	The output from a simulation, as obtained from <code>oncoSimulIndiv</code> , <code>oncoSimulPop</code> , or <code>oncoSimulSample</code> (see <a href="#">oncoSimulIndiv</a> ). This must be from v.2 and forward (no phylogenetic information is stored for earlier objects).
N	Show in the plot all clones that have a population size of at least N at time <code>time</code> and the parents of those clones (parents are shown regardless of population size —i.e., you can see extinct parents). If you want to show everything that ever appeared, set <code>N = 0</code> .
t	The time at which N should be satisfied. This can either be the string "last", meaning the last time of the simulation, or a range of two values. In the second case, all clones with population size of at least N in at least one time point between <code>time[1]</code> and <code>time[2]</code> will be shown (together with their parents).
timeEvents	If TRUE, the vertical position of the nodes in the plot will be proportional to their time of first appearance.
keepEvents	If TRUE, the graph will show all the birth events. Thus, the number of arrows shows the number of times a clone give rise to another. For large graphs with many events, this slows the graph considerably.
fixOverlap	When using <code>timeEvents = TRUE</code> nodes can overlap (as we modify their vertical location after <code>igraph</code> has done the initial layout). This attempts to fix that problem by randomly relocating, along the X axis, the nodes that have the same X value.
returnGraph	If TRUE, the <code>igraph</code> object is returned. You can use this to plot the object however you want or obtain the adjacency matrix.
...	Additional arguments. Currently not used..

**Value**

A plot is produced. If `returnGraph` the `igraph` object is returned.

**Note**

These are not, technically, proper phylogenetic trees and we use "phylogeny" here in an abuse of terminology. The plots we use, where we show parent child relationships are arguably more helpful in this context. But you could draw proper phylogenies with the information provided.

If you want to obtain the adjacency matrix, this is trivial: just set `returnGraph = TRUE` and use [get.adjacency](#). See an example below.

**Author(s)**

Ramon Diaz-Uriarte

**See Also**

[oncoSimulIndiv](#)

**Examples**

```

data(examplesFitnessEffects)
tmp <- oncoSimulIndiv(examplesFitnessEffects[["o3"]],
  model = "McFL",
  mu = 5e-5,
  detectionSize = 1e8,
  detectionDrivers = 3,
  sampleEvery = 0.025,
  max.num.tries = 10,
  keepEvery = 5,
  initSize = 2000,
  finalTime = 3000,
  onlyCancer = FALSE,
  keepPhylog = TRUE)

## Show only those with N > 10 at end
plotClonePhylog(tmp, N = 10)

## Show only those with N > 1 between times 5 and 1000
plotClonePhylog(tmp, N = 1, t = c(5, 1000))

## Show everything, even if terminal nodes are extinct
plotClonePhylog(tmp, N = 0)

## Show time when first appeared
plotClonePhylog(tmp, N = 10, timeEvents = TRUE)

## Not run:
## Show each event
## This can take a few seconds
plotClonePhylog(tmp, N = 10, keepEvents = TRUE)

## End(Not run)

## Adjacency matrix
require(igraph)
get.adjacency(plotClonePhylog(tmp, N = 10, returnGraph = TRUE))

```

---

plotFitnessLandscape *Plot a fitness landscape.*

---

**Description**

Show a plot of a fitness landscape. The plot is modeled after (actually, mostly a blatant copy of) that of MAGELLAN, <http://wwwabi.snv.jussieu.fr/public/Magellan/>.

Note: this is not a plot of the fitnessEffects object; for that, see [plot.fitnessEffects](#).

**Usage**

```
plotFitnessLandscape(x, show_labels = TRUE,
```

```

        col = c("green4", "red", "yellow"),
        lty = c(1, 2, 3),
        use_ggrepel = FALSE,
        log = FALSE, max_num_genotypes = 2000,
        only_accessible = FALSE,
        accessible_th = 0,
        ...)

## S3 method for class 'genotype_fitness_matrix'
plot(x, show_labels = TRUE,
     col = c("green4", "red", "yellow"),
     lty = c(1, 2, 3),
     use_ggrepel = FALSE,
     log = FALSE, max_num_genotypes = 2000,
     only_accessible = FALSE,
     accessible_th = 0,
     ...)

## S3 method for class 'evalAllGenotypes'
plot(x, show_labels = TRUE,
     col = c("green4", "red", "yellow"),
     lty = c(1, 2, 3),
     use_ggrepel = FALSE,
     log = FALSE, max_num_genotypes = 2000,
     only_accessible = FALSE,
     accessible_th = 0,
     ...)

## S3 method for class 'evalAllGenotypesMut'
plot(x, show_labels = TRUE,
     col = c("green4", "red", "yellow"),
     lty = c(1, 2, 3),
     use_ggrepel = FALSE,
     log = FALSE, max_num_genotypes = 2000,
     only_accessible = FALSE,
     accessible_th = 0,
     ...)

```

## Arguments

- x
- One of the following:
- A matrix (or data frame) with  $g + 1$  columns. Each of the first  $g$  columns contains a 1 or a 0 indicating that the gene of that column is mutated or not. Column  $g + 1$  contains the fitness values. This is, for instance, the output you will get from [rfitness](#).
  - A two column data frame. The second column is fitness, and the first column are genotypes, given as a character vector. For instance, a row "A, B" would mean the genotype with both A and B mutated.
  - The output from a call to [evalAllGenotypes](#). Make sure you use `order = FALSE` in that call.
  - The output from a call to [evalAllGenotypesMut](#). Make sure you use `order = FALSE`.

- The output from a call to `allFitnessEffects`.

The first two are the same as the format for the `genotFitness` component in `allFitnessEffects`.

<code>show_labels</code>	If TRUE, show the genotype labels.
<code>col</code>	A three-element vector that gives the colors to use for increase, decreases and no changes in fitness, respectively. The first two colours are also used for peaks and sinks.
<code>lty</code>	A three-element vector that gives the line types to use for increase, decreases and no changes in fitness, respectively.
<code>use_ggrepel</code>	If TRUE, use the <code>ggrepel</code> package to avoid overlap of labels.
<code>log</code>	Log-scale the y axis.
<code>max_num_genotypes</code>	Maximum allowed number of genotypes. For some types of input, we make a call to <code>evalAllGenotypes</code> , and use this as the maximum.
<code>only_accessible</code>	If TRUE, show only accessible paths. A path is considered accessible if, at each mutational step (i.e., with the addition of each mutation) fitness increases by at least <code>accessible_th</code> . If you set <code>only_accessible = TRUE</code> , the number of genotypes displayed can be much smaller than the number of existing genotypes if many of those genotypes are not accessible via any path.
<code>accessible_th</code>	The threshold for the minimal change in fitness at each mutation step (i.e., between successive genotypes) to be used if <code>only_accessible = TRUE</code> .
<code>...</code>	Other arguments passed to <code>plot</code> . Not used for now.

### Value

A fitness landscape plot: a plot showing paths between genotypes and peaks and sinks (local maxima and minima).

### Note

I have copied most of the ideas (and colors, and labels) of this plot from MAGELLAN (<http://www.abi.snv.jussieu.fr/public/Magellan/>) but MAGELLAN has other functionality that is not provided here such as epistasis stats for the landscape, and several visual manipulation options.

One feature of this function that is not available in MAGELLAN is showing genotype labels (i.e., annotated by gene names), which can be helpful if the different genotypes mean something to you.

In addition to the above differences, another difference between this plot and those of MAGELLAN is **how sinks/peaks of more than one genotype are dealt with**. This plot will show as sinks or peaks sets of one or more genotypes that are of identical fitness (and separated by a Hamming distance of one). So a sink or a peak might actually be made of more than one genotype. In MAGELLAN, as far as I can tell, peaks and sinks are always made of a single isolated genotype.

Does this matter? In most realistic cases where not two genotypes can have exactly the same fitness it does not. In some cases, though, it might matter. Are multi-genotype sinks/peaks really sinks/peaks? Arguably yes: suppose genotypes "AB" and "ABC" both have fitness 0, which is minimal among the fitness in the set of genotypes, and genotypes "A" and "ABCD" have fitness 0.1. To go from "A" to "ABCD", if you want to travel through "AB", you have to go through the valley of "AB" and "ABC"; once in "ABC" you can climb up to "ABCD"; and once in "AB" you can move to "ABC" since it has identical fitness to "AB". Mutatis mutandis for multi-genotype peaks. Ignoring the possibility of peaks/sinks made of more than one genotype actually makes code much simpler.

Sometimes not showing the any links that involve a decrease in fitness can help see non-accessible pathways (in strong selection, no multiple mutations, etc); do this by passing, for instance, an NA for the second element of col.

Finally, use common sense: for instance, if you pass a `allFitnessEffects` that specifies for, say, the fitness of a total of 5000 genotypes you'll have to wait a while for the plot to finish.

### Author(s)

Ramon Diaz-Uriarte

### References

MAGELLAN web site: <http://wwwabi.snv.jussieu.fr/public/Magellan/>

Brouillet, S. et al. (2015). MAGELLAN: a tool to explore small fitness landscapes. *bioRxiv*, **31583**. <http://doi.org/10.1101/031583>

### See Also

[allFitnessEffects](#), [evalAllGenotypes](#), [allFitnessEffects](#), [rfitness](#), [plot.fitnessEffects](#)

### Examples

```
## Generate random fitness for four genes-genotypes
## and plot landscape.

r1 <- rfitness(4)
plot(r1)

## Specify fitness in a matrix, and plot it

m5 <- cbind(A = c(0, 1, 0, 1), B = c(0, 0, 1, 1), F = c(1, 2, 3, 5.5))
plotFitnessLandscape(m5)

## Specify fitness with allFitnessEffects, and plot it

fe <- allFitnessEffects(epistasis = c("a : b" = 0.3,
                                     "b : c" = 0.5),
                       noIntGenes = c("e" = 0.1))

plot(evalAllGenotypes(fe, order = FALSE))

## same as
plotFitnessLandscape(evalAllGenotypes(fe, order = FALSE))
```

---

plotPoset	<i>Plot a poset.</i>
-----------	----------------------

---

### Description

Plot a poset. Optionally add a root and change names of nodes.

### Usage

```
plotPoset(x, names = NULL, addroot = FALSE, box = FALSE, ...)
```

### Arguments

x	A poset. A matrix with two columns where, in each row, the first column is the ancestor and the second the descendant. Note that there might be multiple rows with the same ancestor, and multiple rows with the same descendant. See <a href="#">poset</a> .
names	If not NULL, a vector of names for the nodes, with the same length as the total number of nodes in a poset (which need not be the same as the number of rows; see <a href="#">poset</a> ). If <code>addroot = TRUE</code> , then 1 + the number of nodes in the poset.
addroot	Add a "Root" node to the graph?
box	Should the graph be placed inside a box?
...	Additional arguments to plot (actually, <code>plot.graphNEL</code> in the <code>Rgraphviz</code> package).

### Details

The poset is converted to a graphNEL object.

### Value

A plot is produced.

### Author(s)

Ramon Diaz-Uriarte

### See Also

[examplePosets](#), [poset](#)

### Examples

```
data(examplePosets)
plotPoset(examplePosets[["p1101"]])

## If you will be using that poset a lot, maybe simpler if

poset701 <- examplePosets[["p701"]]
plotPoset(poset701, addroot = TRUE)
```

```

## Compare to Pancreatic cancer figure in Gerstung et al., 2011

plotPoset(poset701,
          names = c("KRAS", "SMAD4", "CDNK2A", "TP53",
                   "MLL3", "PXDN", "TGFB2R2"))

## If you want to show Root explicitly do

plotPoset(poset701, addroot = TRUE,
          names = c("Root", "KRAS", "SMAD4", "CDNK2A", "TP53",
                   "MLL3", "PXDN", "TGFB2R2"))

## Of course, names are in the order of nodes, so KRAS is for node 1,
## etc, but the order of entries in the poset does not matter:

poset701b <- poset701[nrow(poset701):1, ]

plotPoset(poset701b,
          names = c("KRAS", "SMAD4", "CDNK2A", "TP53",
                   "MLL3", "PXDN", "TGFB2R2"))

```

---

POM

*Obtain Lines of Descent and Paths of the Maximum and their diversity from simulations.*

---

## Description

Compute Lines of Descent (LOD) and Path of the Maximum (POM) for a single simulation or a set of simulations (from `oncoSimulPop`).

`diversityPOM` and `diversityLOD` return the Shannon's diversity (entropy) of the POM and LOD, respectively, of a set of simulations (it makes no sense to compute those from a single simulation).

## Usage

```

POM(x)
LOD(x)
diversityPOM(lpom)
diversityLOD(llod)

```

## Arguments

<code>x</code>	An object of class <code>oncosimulpop</code> (version $\geq 2$ , so simulations with the old <code>poset</code> specification will not work) or class <code>oncosimul2</code> (a single simulation).
<code>lpom</code>	A list of POMs, as returned from <code>POM</code> on an object of class <code>oncosimulpop</code> .
<code>llod</code>	A list of LODs, as returned from <code>LOD</code> on an object of class <code>oncosimulpop</code> .
<code>...</code>	Other arguments passed to methods (ignored now).



```

child = c("KRAS", "SMAD4", "CDNK2A",
          "TP53", "MLL3",
          rep("PXDN", 3), rep("TGFB2", 2)),
s = 0.05,
sh = -0.3,
typeDep = "MN"))

pancr1 <- oncoSimulIndiv(pancr, model = "Exp")
pancr8 <- oncoSimulPop(8, pancr, model = "Exp",
                      mc.cores = 2)

POM(pancr1)
LOD(pancr1)

POM(pancr8)
LOD(pancr8)

diversityPOM(POM(pancr8))
diversityLOD(LOD(pancr8))

```

---

poset

*Poset*

---

## Description

Poset: explanation.

## Arguments

x                    The poset. See details.

## Details

A poset is a two column matrix. In each row, the first column is the ancestor (or the restriction) and the second column the descendant (or the node that depends on the restriction). Each node is identified by a positive integer. The graph includes all nodes with integers between 1 and the largest integer in the poset.

Each node can be necessary for several nodes: in this case, the same node would appear in the first column in several rows.

A node can depend on two or more nodes (conjunctions): in this case, the same node would appear in the second column in several rows.

There can be nodes that do not depend on anything (except the Root node) and on which no other nodes depend. The simplest and safest way to deal with all possible cases, including these cases, is to have all nodes with at least one entry in the poset, and nodes that depend on no one, and on which no one depends should be placed on the second column (with a 0 on the first column).

Alternatively, any node not named explicitly in the poset, but with a number smaller than the largest number in the poset, is taken to be a node that depends on no one and on which no one depends. See examples below.

This specification of restrictions is for version 1. See [allFitnessEffects](#) for a much more flexible one for version 2. Both can be used with [oncoSimulIndiv](#).

### Author(s)

Ramon Diaz-Uriarte

### References

Posets and similar structures appear in several places. The following two papers use them extensively.

Gerstung et al., 2009. Quantifying cancer progression with conjunctive Bayesian networks. *Bioinformatics*, 21: 2809–2815.

Gerstung et al., 2011. The Temporal Order of Genetic and Pathway Alterations in Tumorigenesis. *PLoS ONE*, 6.

### See Also

[examplePosets](#), [plotPoset](#), [oncoSimulIndiv](#)

### Examples

```
## Node 2 and 3 depend on 1, and 4 depends on no one
p1 <- cbind(c(1L, 1L, 0L), c(2L, 3L, 4L))
plotPoset(p1, addroot = TRUE)

## Node 2 and 3 depend on 1, and 4 to 7 depend on no one.
## We do not have nodes 4 to 6 explicitly in the poset.
p2 <- cbind(c(1L, 1L, 0L), c(2L, 3L, 7L))
plotPoset(p2, addroot = TRUE)

## But this is arguably cleaner
p3 <- cbind(c(1L, 1L, rep(0L, 4)), c(2L, 3L, 4:7 ))
plotPoset(p3, addroot = TRUE)

## A simple way to create a poset where no gene (in a set of 15) depends
## on any other.

p4 <- cbind(0L, 15L)
plotPoset(p4, addroot = TRUE)

## Specifying the pancreatic cancer poset in Gerstung et al., 2011
## (their figure 2B, left). We use numbers, but for nicer plotting we
## will use names: KRAS is 1, SMAD4 is 2, etc.

pancreaticCancerPoset <- cbind(c(1, 1, 1, 1, 2, 3, 4, 4, 5),
                             c(2, 3, 4, 5, 6, 6, 6, 7, 7))
storage.mode(pancreaticCancerPoset) <- "integer"

plotPoset(pancreaticCancerPoset,
          names = c("KRAS", "SMAD4", "CDNK2A", "TP53",
                   "MLL3", "PXDN", "TGFBR2"))
```

```
## Specifying poset 2 in Figure 2A of Gerstung et al., 2009:

poset2 <- cbind(c(1, 1, 3, 3, 3, 7, 7, 8, 9, 10),
               c(2, 3, 4, 5, 6, 8, 9, 10, 10, 11))

storage.mode(poset2) <- "integer"
plotPoset(poset2)
```

---

rfitness

*Generate random fitness.*


---

### Description

Generate random fitness landscapes under a House of Cards, Rough Mount Fuji, or additive model.

### Usage

```
rfitness(g, c = 0.5, sd = 1, mu = 1, reference = "random", scale = NULL,
         wt_is_1 = c("subtract", "divide", "force", "no"),
         log = FALSE, min_accessible_genotypes = NULL,
         accessible_th = 0, truncate_at_0 = TRUE)
```

### Arguments

<code>g</code>	Number of genes.
<code>c</code>	The decrease in fitness of a genotype per each unit increase in Hamming distance from the reference genotype (see reference).
<code>sd</code>	The standard deviation of the random component (a normal distribution of mean <code>mu</code> and standard deviation <code>sd</code> ).
<code>mu</code>	The mean of the random component (a normal distribution of mean <code>mu</code> and standard deviation <code>sd</code> ).
<code>reference</code>	The reference genotype: for the deterministic, additive part, this is the genotype with maximal fitness, and all other genotypes decrease their fitness by <code>c</code> for every unit of Hamming distance from this reference. If "random" a genotype will be randomly chosen as the reference. If "max" the genotype with all positions mutated will be chosen as the reference. If you pass a vector (e.g., <code>reference = c(1, 0, 1, 0)</code> ) that will be the reference genotype. If "random2" a genotype will be randomly chosen as the reference. In contrast to "random", however, not all genotypes have the same probability of being chosen; here, what is equal is the probability that the reference genotype has 1, 2, ..., <code>g</code> , mutations (and, once a number mutations is chosen, all genotypes with that number of mutations have equal probability of being the reference).
<code>scale</code>	Either NULL (nothing is done) or a two-element vector. If a two-element vector, fitness is re-scaled between <code>scale[1]</code> (the minimum) and <code>scale[2]</code> (the maximum).

wt_is_1	<p>If "divide" (the default) the fitness of all genotypes is divided by the fitness of the wildtype (after possibly adding a value to ensure no negative fitness) so that the wildtype (the genotype with no mutations) has fitness 1. This is a case of scaling, and it is applied after scale, so if you specify both "wt_is_1 = 'divide'" and use an argument for scale it is most likely that the final fitness will not respect the limits in scale.</p> <p>If "subtract" we shift all the fitness values (subtracting fitness of the wildtype and adding 1) so that the wildtype ends up with a fitness of 1. This is also applied after scale, so if you specify both "wt_is_1 = 'subtract'" and use an argument for scale it is most likely that the final fitness will not respect the limits in scale (though the distortion might be simpler to see as just a shift up or down).</p> <p>If "force" we simply set the fitness of the wildtype to 1, without any divisions. This means that the scale argument would work (but it is up to you to make sure that the range of the scale argument includes 1 or you might not get what you want). Note that using this option can easily lead to landscapes with no accessible genotypes (even if you also use scale).</p> <p>If "none", the fitness of the wildtype is not touched.</p>
log	If TRUE, log-transform fitness.
min_accessible_genotypes	<p>If not NULL, the minimum number of accessible genotypes in the fitness landscape. A genotype is considered accessible if you can reach it from the wildtype by going through at least one path where all changes in fitness are larger or equal to <code>accessible_th</code>. The changes in fitness are considered at each mutational step, i.e., at each addition of one mutation we compute the difference between the genotype with <math>k + 1</math> mutations minus the ancestor genotype with <math>k</math> mutations. Thus, a genotype is considered accessible if there is at least one path where fitness increases at each mutational step by at least <code>accessible_th</code>.</p> <p>If the condition is not satisfied, we continue generating random fitness landscapes with the specified parameters until the condition is satisfied.</p> <p>(Why check against NULL and not against zero? Because this allows you to count accessible genotypes even if you do not want to ensure a minimum number of accessible genotypes.)</p>
accessible_th	The threshold for the minimal change in fitness at each mutation step (i.e., between successive genotypes) that allows a genotype to be regarded as accessible. This only applies if <code>min_accessible_genotypes</code> is larger than 0. So if you want to allow small decreases in fitness in successive steps, use a small negative value for <code>accessible_th</code> .
truncate_at_0	If TRUE (the default) any fitness $\leq 0$ is substituted by a small positive constant ( $1e-9$ ). Why? Because MAGELLAN and some plotting routines can have trouble (specially if you log) with values $\leq 0$ . Or we might have trouble if we want to log the fitness.

## Details

The model used here follows the Rough Mount Fuji model in Szendro et al., 2013 or Franke et al., 2011. Fitness is given as

$$f(i) = -cd(i, reference) + x_i$$

where  $d(i, j)$  is the Hamming distance between genotypes  $i$  and  $j$  (the number of positions that differ) and  $x_i$  is a random variable (in this case, a normal deviate of mean  $\mu$  and standard deviation  $sd$ ).

Setting  $c = 0$  we obtain a House of Cards model. Setting  $sd = 0$  fitness is given by the distance from the reference and if the reference is the genotype with all positions mutated, then we have a fully additive model (fitness increases linearly with the number of positions mutated).

For OncoSimulR, we often want the wildtype to have a mean of 1. Reasonable settings are  $\mu = 1$  and  $wt\_is\_1 = 'subtract'$  so that we simulate from a distribution centered in 1, and we make sure afterwards (via a simple shift) that the wildtype is actual 1. The  $sd$  controls the standard deviation, with the usual working and meaning as in a normal distribution, unless  $c$  is different from zero. In this case, with  $c$  large, the range of the data can be large, specially if  $g$  (the number of genes) is large.

### Value

An matrix with  $g + 1$  columns. Each column corresponds to a gene, except the last one that corresponds to fitness. 1/0 in a gene column denotes gene mutated/not-mutated. (For ease of use in other functions, this matrix has class "genotype\_fitness\_matrix".)

If you have specified `min_accessible_genotypes > 0`, the return object has added attributes `accessible_genotypes` and `accessible_th` that show the number of accessible genotypes under the specified threshold.

### Author(s)

Ramon Diaz-Uriarte

### References

Szendro I.-G. et al. (2013). Quantitative analyses of empirical fitness landscapes. *Journal of Statistical Mechanics: Theory and Experiment*, **01**, P01005.

Franke, J. et al. (2011). Evolutionary accessibility of mutational pathways. *PLoS Computational Biology*, **7**(8), 1–9.

### See Also

[oncoSimulIndiv](#), [plot.genotype\\_fitness\\_matrix](#), [evalAllGenotypes](#) [allFitnessEffects](#) [plotFitnessLandscape](#)

### Examples

```
## Random fitness for four genes-genotypes,
## plotting and simulating an oncogenetic trajectory

r1 <- rfittness(4)
plot(r1)
oncoSimulIndiv(allFitnessEffects(genotFitness = r1))
```

---

<code>samplePop</code>	<i>Obtain a sample from a population of simulations.</i>
------------------------	--

---

### Description

Obtain a sample (a matrix of individuals/samples by genes or, equivalently, a vector of "genotypes") from an `oncosimulpop` object (i.e., a simulation of multiple individuals) or a single `oncosimul` object. Sampling schemes include whole tumor and single cell sampling, and sampling at the end of the tumor progression or during the progression of the disease.

`sampledGenotypes` shows the genotype frequencies from that sample; Shannon's diversity — entropy— of the genotypes is also returned. Order effects are ignored.

### Usage

```
samplePop(x, timeSample = "last", typeSample = "whole",
          thresholdWhole = 0.5, geneNames = NULL, popSizeSample = NULL,
          propError = 0)
```

```
sampledGenotypes(y, genes = NULL)
```

### Arguments

<code>x</code>	An object of class <code>oncosimulpop</code> or class <code>oncosimul2</code> (a single simulation).
<code>y</code>	The output from a call to <code>samplePop</code> .
<code>timeSample</code>	"last" means to sample each individual in the very last time period of the simulation. "unif" (or "uniform") means sampling each individual at a time chosen uniformly from all the times recorded in the simulation between the time when the first driver appeared and the final time period. "unif" means that it is almost sure that different individuals will be sampled at different times. "last" does not guarantee that different individuals will be sampled at the same time unit, only that all will be sampled in the last time unit of their simulation. You can, alternatively, specify the population size at which you want the sample to be taken. See argument <code>popSizeSample</code> .
<code>typeSample</code>	"singleCell" (or "single") for single cell sampling, where the probability of sampling a cell (a clone) is directly proportional to its population size. "wholeTumor" (or "whole") for whole tumor sampling (i.e., this is similar to a biopsy being the entire tumor). "singleCell-noWT" or "single-nowt" is single cell sampling, but excluding the wild type.
<code>thresholdWhole</code>	In whole tumor sampling, whether a gene is detected as mutated depends on <code>thresholdWhole</code> : a gene is considered mutated if it is altered in at least <code>thresholdWhole</code> proportion of the cells in that individual.
<code>geneNames</code>	An optional vector of gene names so as to label the column names of the output.
<code>popSizeSample</code>	An optional vector of total population sizes at which you want the samples to be taken. If you pass this vector, <code>timeSample</code> has no effect. The samples will be taken at the first time at which the population size gets as large as (or larger than) the size specified in <code>popSizeSample</code> . This allows you to specify arbitrary sampling schemes with respect to total population size.

propError	The proportion of observations with error (for instance, genotyping error). If larger than 0, this proportion of entries in the sampled matrix will be flipped (i.e., 0s turned to 1s and 1s turned to 0s).
genes	If non-NULL, use only the genes in genes to create the table of genotypes. This can be useful if you only care about the genotypes with respect to a subset of genes (say, X), and want to collapse with respect to another subset of genes (say, Y), for instance if Y is a large set of passenger genes. For example, suppose the complete set of genes is 'a', 'b', 'c', 'd', and you specify genes = c('a', 'b'); then, genotypes 'a, b, c' and genotypes 'a, b, d' will not be shown as different rows in the table of frequencies. Likewise, genotypes 'a, c' and genotypes 'a, d' will not be shown as different rows. Of course, if what are actually different genotypes are not regarded as different, this will affect the calculation of the diversity.

## Details

samplePop simply repeats the sampling process in each individual of the oncosimulpop object.

Please see [oncoSimulSample](#) for a much more efficient way of sampling when you are sure what you want to sample.

Note that if you have set `onlyCancer = FALSE` in the call to [oncoSimulSample](#), you can end up trying to sample from simulations where the population size is 0. In this case, you will get a vector/matrix of NAs and a warning.

Similarly, when using `timeSample = "last"` you might end up with a vector of 0 (not NAs) because you are sampling from a population that contains no clones with mutated genes. This event (sampling from a population that contains no clones with mutated genes), by construction, cannot happen when `timeSample = "unif"` as "uniform" sampling is taken here to mean sampling at a time chosen uniformly from all the times recorded in the simulation between the time when the first driver appeared and the final time period. However, you might still get a vector of 0, with uniform sampling, if you sample from a population that contains only a few cells with any mutated genes, and most cells with no mutated genes.

## Value

A matrix. Each row is a "sample genotype", where 0 denotes no alteration and 1 alteration. When using v.2, columns are named with the gene names.

We quote "sample genotype" because when not using single cell, a row (a sample genotype) need not be, of course, any really existing genotype in a population as we are genotyping a whole tumor. Suppose there are really two genotypes present in the population, genotype A, which has gene A mutated and genotype B, which has gene B mutated. Genotype A has a frequency of 60% (so B's frequency is 40%). If you use whole tumor sampling with `thresholdWhole = 0.4` you will obtain a genotype with A and B mutated.

For `sampldGenotypes` a data frame with two columns: `genotypes` and `frequencies`. This data frame has an additional attribute, "ShannonI", where Shannon's index of diversity (entropy) is stored. This is an object of class "sampldGenotypes" with an S3 print method.

## Author(s)

Ramon Diaz-Uriarte

## References

Diaz-Uriarte, R. (2015). Identifying restrictions in the order of accumulation of mutations during tumor progression: effects of passengers, evolutionary models, and sampling <http://www.biomedcentral.com/1471-2105/16/41/abstract>

## See Also

[oncoSimulPop](#), [oncoSimulSample](#)

## Examples

```
data(examplePosets)
p705 <- examplePosets[["p705"]]

## (I set mc.cores = 2 to comply with --as-cran checks, but you
## should either use a reasonable number for your hardware or
## leave it at its default value).

p1 <- oncoSimulPop(4, p705, mc.cores = 2)
(sp1 <- samplePop(p1))
sampledGenotypes(sp1)

## Sample at fixed sizes. Notice the requested size
## for the last population is larger than the any population size
## so we get NAs

(sp2 <- samplePop(p1, popSizeSample = c(1e7, 1e6, 4e5, 1e13)))
sampledGenotypes(sp2)

## Now single cell sampling

r1 <- oncoSimulIndiv(p705)
samplePop(r1, typeSample = "single")

sampledGenotypes(samplePop(r1, typeSample = "single"))
```

---

simOGraph

*Simulate oncogenetic/CBN/XMPN DAGs.*

---

## Description

Simulate DAGs that represent restrictions in the accumulation of mutations.

## Usage

```
simOGraph(n, h = ifelse(n >= 4, 4, n), conjunction = TRUE, nparents = 3,
multilevelParent = TRUE, removedirectIndirect = TRUE, rootName = "Root",
geneNames = seq.int(n), out = c("adjmat", "rT"),
s = 0.1, sh = -0.1, typeDep = "AND")
```

**Arguments**

n	Number of nodes, or edges, in the graph. Like the number of genes.
h	Approximate height of the graph. See details.
conjunction	If TRUE, conjunctions (i.e., multiple parents for a node) are allowed.
nparents	Maximum number of parents of a node, when conjunction is TRUE.
multilevelParent	Can a node have parents at different heights (i.e., parents that are at different distance from the root node)?
removeDirectIndirect	Ensure that no two nodes are connected both directly (i.e., with an edge between them) and indirectly, through intermediate nodes. If TRUE, we return the transitive reduction of the DAG.
rootName	The name you want to give the "Root" node.
geneNames	The names you want to give the the non-root nodes.
out	Whether the ouptut should be an adjacency matrix or a "restriction table", as used in <a href="#">allFitnessEffects</a> .
s	If using as output a restriction, the default value for s. See <a href="#">allFitnessEffects</a> .
sh	If using as output a restriction, the default value for sh. See <a href="#">allFitnessEffects</a>
typeDep	If using as output a restriction, the default value for "typeDep". See <a href="#">allFitnessEffects</a>

**Details**

This is a simple, heuristic procedure for generating graphs of restrictions that seem compatible with published trees in the oncogenetic literature.

The basic procedure is as follows: nodes (argument *n*) are split into approximately equally sized *h* groups, and then each node from a level is connected to nodes chosen randomly from nodes of the remaining superior (i.e., closer to the Root) levels. The number of edges comes from a uniform distribution between 1 and *nparents*.

The actual depth of the graph can be smaller than *h* because nodes from a level might be connected to superior levels skipping intermediate ones.

See the vignette for further discussion about arguments.

**Value**

An adjacency matrix for a directed graph or a data frame to be used as input, as "restriction table" in [allFitnessEffects](#).

**Author(s)**

Ramon Diaz-Uriarte

**Examples**

```
(a1 <- simOGraph(10))
library(graph) ## for simple plotting
plot(as(a1, "graphNEL"))

simOGraph(3, geneNames = LETTERS[1:3])
```

---

to_Magellan	<i>Create output for MAGELLAN.</i>
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---

### Description

Create a fitness landscape in a format that is understood by MAGELLAN <http://wwwabi.snv.jussieu.fr/public/Magellan/>.

### Usage

```
to_Magellan(x, file,  
            max_num_genotypes = 2000)
```

### Arguments

x	One of the following: <ul style="list-style-type: none"><li>• A matrix (or data frame) with <math>g + 1</math> columns. Each of the first <math>g</math> columns contains a 1 or a 0 indicating that the gene of that column is mutated or not. Column <math>g + 1</math> contains the fitness values. This is, for instance, the output you will get from <a href="#">rfitness</a>.</li><li>• A two column data frame. The second column is fitness, and the first column are genotypes, given as a character vector. For instance, a row "A, B" would mean the genotype with both A and B mutated.</li><li>• The output from a call to <a href="#">evalAllGenotypes</a>. Make sure you use <code>order = FALSE</code> in that call.</li><li>• The output from a call to <a href="#">evalAllGenotypesMut</a>. Make sure you use <code>order = FALSE</code>.</li><li>• The output from a call to <a href="#">allFitnessEffects</a> (with no order effects in the specification).</li></ul> The first two are the same as the format for the <code>genotFitness</code> component in <a href="#">allFitnessEffects</a> .
file	The name of the output file. If NULL, a name will be created using <a href="#">tempfile</a> .
max_num_genotypes	Maximum allowed number of genotypes. For some types of input, we make a call to <a href="#">evalAllGenotypes</a> , and use this as the maximum.

### Value

A file is written to disk. You can then plot and/or show summary statistics using MAGELLAN.

### Note

If you try to pass a fitness specification with order effects you will receive an error, since that cannot be plotted with MAGELLAN.

### Author(s)

Ramon Diaz-Uriarte

## References

MAGELLAN web site: <http://wwwabi.snv.jussieu.fr/public/Magellan/>

Brouillet, S. et al. (2015). MAGELLAN: a tool to explore small fitness landscapes. *bioRxiv*, **31583**.  
<http://doi.org/10.1101/031583>

## See Also

[allFitnessEffects](#), [evalAllGenotypes](#), [allFitnessEffects](#), [rfitness](#)

## Examples

```
## Generate random fitness for four-genes genotype
## and export landscape.

r1 <- rfittness(4)
to_Magellan(r1, NULL)

## Specify fitness using a DAG and export it
cs <- data.frame(parent = c(rep("Root", 3), "a", "d", "c"),
                 child = c("a", "b", "d", "e", "c", "f"),
                 s = 0.1,
                 sh = -0.9,
                 typeDep = "MN")

to_Magellan(allFitnessEffects(cs), NULL)
```

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