# OncoSimulR: forward genetic simulation in asexual populations with arbitrary epistatic interactions and a focus on modeling tumor progression.

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# 1 Introduction

OncoSimulR is an individual- or clone-based forward-time genetic simulator for biallelic markers (wildtype vs. mutated) in asexually reproducing populations without spatial structure (perfect mixing). Its design emphasizes flexible specification of fitness and mutator effects.

OncoSimulR was originally developed to simulate tumor progression with emphasis on allowing users to set restrictions in the accumulation of mutations as specified, for example, by Oncogenetic Trees (OT: Desper et al. 1999; Szabo and Boucher 2008) or Conjunctive Bayesian Networks (CBN: Beerenwinkel, Eriksson, and Sturmfels 2007; Gerstung et al. 2009, 2011), with the possibility of adding passenger mutations to the simulations and allowing for several types of sampling.

Since then, OncoSimulR has been vastly extended to allow you to specify other types of restrictions in the accumulation of genes, such as the XOR models of Korsunsky et al. (2014) or the "semimonotone" model of Farahani and Lagergren (2013). Moreover, different fitness effects related to the order in which mutations appear can also be incorporated, involving arbitrary numbers of genes. This is *very* different from "restrictions in the order of accumulation of mutations". With order effects, described in a recent cancer paper by Ortmann and collaborators (Ortmann et al. 2015), the effect of having both mutations "A" and "B" differs depending on whether "A" appeared before or after "B" (the actual case involves genes JAK2 and TET2).

More generally, OncoSimulR now also allows you to specify arbitrary epistatic interactions between arbitrary collections of genes and to model, for example, synthetic mortality or synthetic viability (again, involving an arbitrary number of genes, some of which might also depend on other genes, or show order effects with other genes). Moreover, it is possible to specify the above interactions in terms of modules, not genes. This idea is discussed in, for example, Raphael and Vandin (2015) and Gerstung et al. (2011): the restrictions encoded in, say, CBNs or OT can be considered to apply not to genes, but to modules, where each module is a set of genes (and the intersection between modules is the empty set) that performs a specific biological function. Modules, then, play the role of a "union operation" over the set of genes in a module. In addition, arbitrary numbers of genes without interactions (and with fitness effects coming from any distribution you might want) are also possible.

You can also directly specify the mapping between genotypes and fitness and, thus, you can simulate on fitness landscapes of arbitrary complexity.

It is now (released initially in this repo as the freq-dep-fitness branch on February 2019) also possible to simulate scenarios with frequency-dependent fitness, where the fitness of one or more genotypes depends on the relative or absolute frequencies of other genotypes, as in game theory and adaptive dynamics. This makes it possible to model predation and parasitism, cooperation and mutualism, and commensalism.

Mutator/antimutator genes, genes that alter the mutation rate of other genes (Gerrish et al. 2007; Tomlinson, Novelli, and Bodmer 1996), can also be simulated with OncoSimulR and specified with most of the mechanisms above (you can have, for

instance, interactions between mutator genes). And, regardless of the presence or not of other mutator/antimutator genes, different genes can have different mutation rates.

Simulations can be stopped as a function of total population size, number of mutated driver genes, or number of time periods. Simulations can also be stopped with a stochastic detection mechanism where the probability of detecting a tumor increases with total population size. Simulations return the number of cells of every genotype/clone at each of the sampling periods and we can take samples from the former with single-cell or whole- tumor resolution, adding noise if we want. If we ask for them, simulations also store and return the genealogical relationships of all clones generated during the simulation.

The models so far implemented are all continuous time models, which are simulated using the BNB algorithm of Mather, Hasty, and Tsimring (2012). The core of the code is implemented in C++, providing for fast execution. To help with simulation studies, code to simulate random graphs of the kind often seen in CBNs, OTs, etc, is also available. Finally, OncoSimulR also allows for the generation of random fitness landscapes and the representation of fitness landscapes and provides statistics of evolutionary predictability.

## 1.1 Key features of OncoSimulR

As mentioned above, OncoSimulR is now a very general package for forward genetic simulation, with applicability well beyond tumor progression. This is a summary of some of its key features:

- You can specify arbitrary interactions between genes, with arbitrary fitness effects, with explicit support for:
  - Restrictions in the accumulations of mutations, as specified by Oncogenetic Trees (OTs), Conjunctive Bayesian Networks (CBNs), semimonotone progression networks, and XOR relationships.
  - Epistatic interactions including, but not limited to, synthetic viability and synthetic lethality.
  - Order effects.
- You can add passenger mutations.
- You can add mutator/antimutator effects.
- Fitness and mutation rates can be gene-specific.
- You can add arbitrary numbers of non-interacting genes with arbitrary fitness effects.
- you can allow for deviations from the OT, CBN, semimonotone, and XOR models, specifying a penalty for such deviations (the  $s_h$  parameter).

- You can conduct multiple simulations, and sample from them with different temporal schemes and using both whole tumor or single cell sampling.
- You can stop the simulations using a flexible combination of conditions: final time, number of drivers, population size, fixation of certain genotypes, and a stochastic stopping mechanism that depends on population size.
- Right now, three different models are available, two that lead to exponential growth, one of them loosely based on Bozic et al. (2010), and another that leads to logistic-like growth, based on McFarland et al. (2013).
- You can use large numbers of genes (e.g., see an example of 50000 in section 6.5.3).
- Simulations are generally very fast: I use C++ to implement the BNB algorithm (see sections 14.5 and 14.6 for more detailed comments on the usage of this algorithm).
- You can obtain the true sequence of events and the phylogenetic relationships between clones (see section 14.1 for the details of what we mean by "clone").
- You can generate random fitness landscapes (under the House of Cards, Rough Mount Fuji, or additive models, or combinations of the former and under the NK model) and use those landscapes as input to the simulation functions.
- You can plot fitness landscapes.
- You can obtain statistics of evolutionary predictability from the simulations.
- You can now also use simulations with frequency-dependent fitness: fitness (birth rate) is not fixed for a genotype, but can be a function of the frequecies of the clones (see section 10). We can therefore use OncoSimulR to examine, via simulations, results from game theory and adaptive dynamics and study complex scenarios that are not amenable to analytical solutions. More generally, we can model predation and parasitism, cooperation and mutualism, and commensalism.

The table below, modified from the table at the Genetics Simulation Resources (GSR) page, provides a summary of the key features of OncoSimulR. (An explanation of the meaning of terms specific to the GSR table is available from https://popmodels.cancercontrol.cancer.gov/gsr/search/ or from the Genetics Simulation Resources table itself, by moving the mouse over each term).

Table 1: Key features of OncoSimulR. Modified from the original table from https://popmodels.cancercontrol.cancer.gov/gsr/packages/oncosimulr/#detailed .

Attribute Category	Attribute			
Target				
Type of Simulated Data	Haploid DNA Sequence			
Variations	Biallelic Marker, Genotype or Sequencing Error			
Simulation Method	Forward-time			
Type of Dynamical Model	Continuous time			
Entities Tracked	Clones (see 14.2)			
Input	Program specific (R data frames and matrices specifying genotypes' fitness, gene effects, and starting genotype)			
Output				
Data Type	Genotype or Sequence, Individual Relationship (complete parent-child relationships between clones), Demographic (populations sizes of all clones at sampling times), Diversity Measures (LOD, POM, diversity of genotypes), Fitness			
Sample Type	Random or Independent, Longitudinal, Other (proportional to population size)			
Evolutionary Features				
Mating Scheme	Asexual Reproduction			
Demographic				
Population Size Changes	Exponential (two models), Logistic (McFarland et al., 2013)			
Fitness Components				
Birth Rate	Individually Determined from Genotype (models "Exp", "McFL", "McFLD"). Frequency-Dependently Determined from Genotype (models "Exp", "McFL", "McFLD")			
Death Rate	Individually Determined from Genotype (mode "Bozic"), Influenced by Environment—population size (models "McFL" and "McFLD")			
Natural Selection				
Determinant	Single and Multi-locus, Fitness of Offspring, Environmental Factors (population size, genotype frequencies)			
Models	Directional Selection, Multi-locus models, Epistasis, Random Fitness Effects, Frequency-Dependent			
Mutation Models	Two-allele Mutation Model (wildtype, mutant) without back mutation			

Attribute Category	Attribute			
Events Allowed	Varying Genetic Features: change of individual mutation rates (mutator/antimutator genes)			
Spatial Structure	No Spatial Structure (perfectly mixed and no migration)			

Further details about the original motivation for wanting to simulate data this way in the context of tumor progression can be found in Diaz-Uriarte (2015), where additional comments about model parameters and caveats are discussed.

Are there similar programs? The Java program by Reiter et al. (2013), TTP, offers somewhat similar functionality to the previous version of OncoSimulR, but it is restricted to at most four drivers (whereas v.1 of OncoSimulR allowed for up to 64), you cannot use arbitrary CBNs or OTs (or XORs or semimonotone graphs) to specify restrictions, there is no allowance for passengers, and a single type of model (a discrete time Galton-Watson process) is implemented. The current functionality of OncoSimulR goes well beyond the previous version (and, thus, also the TPT of Reiter et al. (2013)). We now allow you to specify all types of fitness effects in other general forward genetic simulators such as FFPopSim (Zanini and Neher 2012), and some that, to our knowledge (e.g., order effects) are not available from any genetics simulator. In addition, the "Lego system" to flexibly combine different fitness specifications is also unique; by "Lego system" I mean that we can combine different pieces and blocks, similarly to what we do with Lego bricks. (I find this an intuitive and very graphical analogy, which I have copied from Hothorn et al. (2006) and Hothorn et al. (2008)). In a nutshell, salient features of OncoSimulR compared to other simulators are the unparalleled flexibility to specify fitness and mutator effects, with modules and order effects as particularly unique, and the options for sampling and stopping the simulations, particularly convenient in cancer evolution models. Also unique in this type of software is the addition of functions for simulating fitness landscapes and assessing evolutionary predictability.

# 1.2 What kinds of questions is OncoSimulR suited for?

OncoSimulR can be used to address questions that span from the effect of mutator genes in cancer to the interplay between fitness landscapes and mutation rates. The main types of questions that OncoSimulR can help address involve combinations of:

- Simulating asexual evolution (the oncoSimul\* functions) where:
  - Fitness is:
    - \* A function of specific epistatic effects between genes
    - \* A function of order effects
    - \* A function of epistatic effects specified using DAGs/posets where these DAGs/posets:
      - · Are user-specified
      - · Generated randomly (simOGraph)

- \* Any mapping between genotypes and fitness where this mapping is:
  - · User-specified
  - · Generated randomly from families of random fitness landscapes (rfitness)
- \* A function of the frequency of other genotypes (i.e., frequency-dependent fitness), such as in adaptive dynamics (see section 10 for more details). This also allows you to model competition, cooperation and mutualism, parasitism and predation, and commensalism between clones.
- Mutation rates can:
  - \* Vary between genes
  - \* Be affected by other genes
- Examining times to evolutionarily or biomedically relevant events (fixation of genotypes, reaching a minimal size, acquiring a minimal number of driver genes, etc—specified with the stopping conditions to the oncoSimul\* functions).
- Using different sampling schemes (samplePop) that are related to:
  - Assessing genotypes from single-cell vs. whole tumor (or whole population)
     with the typeSample argument
  - Genotyping error (propError argument)
  - Timing of samples (timeSample argument)
  - and assessing the consequences of those on the observed genotypes and their diversity (sampledGenotypes) and any other inferences that depend on the observational process.
  - (OncoSimulR returns the abundances of all genotypes at each of the sampling points, so you are not restricted by what the samplePop function provides.)
- Tracking the genealogical relationships of clones (plotClonePhylog) and assessing evolutionary predictability (LOD, POM).

Some specific questions that you can address with the help of OncoSimulR are discussed in section 1.3.

A quick overview of the main functions and their relationships is shown in Figure 1, where we use italics for the type/class of R object and courier font for the name of the functions.

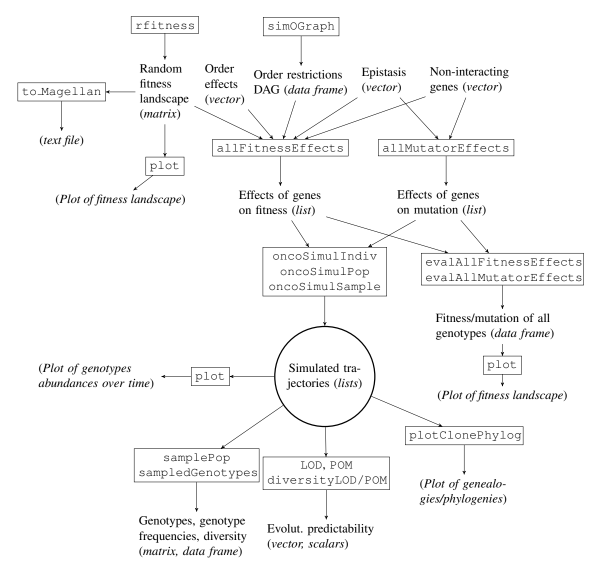


Figure 1: Relationships between the main functions in OncoSimulR.

# 1.3 Examples of questions that can be addressed with OncoSimulR

Most of the examples in the rest of this vignette, starting with those in 1.6, focus on the mechanics. Here, we will illustrate some problems in cancer genomics and evolutionary genetics where OncoSimulR could be of help. This section does not try to provide an answer to any of these questions (those would be full papers by themselves). Instead, this section simply tries to illustrate some kinds of questions where you can use OncoSimulR; of course, the possible uses of OncoSimulR are only limited by your ingenuity. Here, I will only use short snippets of working code as we are limited by time of execution; for real work you would want to use many more scenarios and many more simulations, you would use appropriate statistical methods to compare the output of runs, etc, etc, etc.

```
## Load the package
library(OncoSimulR)
```

#### 1.3.1 Recovering restrictions in the order of accumulation of mutations

This is a question that was addressed, for instance, in Diaz-Uriarte (2015): do methods that try to infer restrictions in the order of accumulation of mutations (e.g., Szabo and Boucher 2008; Gerstung et al. 2009; Ramazzotti et al. 2015) work well under different evolutionary models and with different sampling schemes? (This issue is also touched upon in section 15.2).

A possible way to examine that question would involve:

- generating random DAGs that encode restrictions;
- simulating cancer evolution using those DAGs;
- sampling the data and adding different levels of noise to the sampled data;
- running the inferential method;
- comparing the inferred DAG with the original, true, one.

```
## You would now run the appropriate inferential method and
## compare observed and true. For example

## require(Oncotree)
## fit1 <- oncotree.fit(d1)

## Now, you'd compare fitted and original. This is well beyond
## the scope of this document (and OncoSimulR itself).</pre>
```

#### 1.3.2 Sign epistasis and probability of crossing fitness valleys

This question, and the question in the next section (1.3.3), encompass a wide range of issues that have been addressed in evolutionary genetics studies and which include from detailed analysis of simple models with a few uphill paths and valleys as in Weissman et al. (2009) or Ochs and Desai (2015), to questions that refer to larger, more complex fitness landscapes as in Szendro, Franke, et al. (2013) or Franke et al. (2011) (see below).

Using as an example Ochs and Desai (2015) (we will see this example again in section 5.3, where we cover different ways of specifying fitness), we could specify the fitness landscape and run simulations until fixation (with argument fixation to oncoSimulPop—see more details in section 6.3.3 and 6.3.4, again with this example). We would then examine the proportion of genotypes fixed under different scenarios. And we can extend this example by adding mutator genes:

```
odm <- allMutatorEffects(noIntGenes = c("i" = 50))</pre>
## How do mutation and fitness look like for each genotype?
evalAllGenotypesFitAndMut(od, odm, addwt = TRUE)
     Genotype Fitness MutatorFactor
## 1
           WT
                1.000
## 2
            i
                 0.980
                                   50
                1.200
## 3
            u
                                    1
## 4
                0.000
                                    1
            v
## 5
         i, u
                0.000
                                   50
## 6
         i. v
                1.568
                                   50
## 7
         u, v
                 0.000
                                    1
                                   50
## 8 i, u, v
                 0.000
```

Ochs and Desai explicitly say "Each simulated population was evolved until either the uphill genotype or valley-crossing genotype fixed." So we will use fixation.

```
## Set a small initSize, as o.w. unlikely to pass the valley
initS <- 10
## The number of replicates is tiny, 10, for the sake of speed
## of creation of the vignette
od sim <- oncoSimulPop(10, od, muEF = odm,
                       fixation = c("u", "i, v"), initSize = initS,
                       model = "McFL",
                       mu = 1e-4, detectionDrivers = NA,
                       finalTime = NA,
                       detectionSize = NA, detectionProb = NA,
                       onlyCancer = TRUE,
                       mc.cores = 2, ## adapt to your hardware
                       seed = NULL) ## for reproducibility
## What is the frequency of each final genotype?
sampledGenotypes(samplePop(od sim))
##
##
   Subjects by Genes matrix of 10 subjects and 3 genes.
##
     Genotype Freq
## 1
         i, v
                 4
## 2
                 6
##
##
   Shannon's diversity (entropy) of sampled genotypes: 0.673
```

#### 1.3.3 Predictability of evolution in complex fitness landscapes

Focusing now on predictability in more general fitness landscapes, we would run simulations under random fitness landscapes with varied ruggedness, and would then examine the evolutionary predictability of the trajectories with measures such as "Lines of Descent" and "Path of the Maximum" (Szendro, Franke, et al. 2013) and

the diversity of the sampled genotypes under different sampling regimes (see details in section 12).

```
## For reproducibility
set.seed(7)
RNGkind("L'Ecuyer-CMRG")
## Repeat the following loop for different combinations of whatever
## interests you, such as number of genes, or distribution of the
## c and sd (which affect how rugged the landscape is), or
## reference genotype, or evolutionary model, or stopping criterion,
## or sampling procedure, or ...
## Generate a random fitness landscape, from the Rough Mount
## Fuji model, with g genes, and c ("slope" constant) and
## reference chosen randomly (reference is random by default and
## thus not specified below). Require a minimal number of
## accessible genotypes
g <- 6
c \leftarrow runif(1, 1/5, 5)
rl <- rfitness(g, c = c, min_accessible_genotypes = g)</pre>
## Plot it if you want; commented here as it takes long for a
## viqnette
## plot(rl)
## Obtain landscape measures from MAGELLAN. Export to MAGELLAN and
## call your own copy of MAGELLAN's binary
to_Magellan(rl, file = "rl1.txt")
## or use the binary copy provided with OncoSimulR
## see also below.
Magellan stats(rl)
##
        ngeno
                 npeaks
                             nsinks
                                         qamma
                                                   gamma.
                                                                  r.s
##
       64.000
                  2.000
                              1.000
                                         0.769
                                                    0.854
                                                                0.372
                              nori
##
     nchains
                nsteps
                                         depth
                                                     magn
                                                                sign
##
        1.000
                 5.000
                              4.000
                                        2.000
                                                    0.863
                                                                0.129
##
                   f.1.
                               X.2.
                                         f.3..
                                                               outD m
        rsign
                                                   mode_f
                              0.010
##
       0.008
                  0.916
                                         0.074
                                                    1.000
                                                               0.346
##
       outD_v
                            reach m
                                        fitG_m
                                                    opt_i mProbOpt_0
                 steps\_m
##
       1.587
                   3.052
                             13.597
                                        32.452
                                                    60.000
                                                                0.128
      opt_i.1 mProbOpt_1
##
##
      63.000
                   0.872
```

```
## Simulate evolution in that landscape many times (here just 10)
simulrl <- oncoSimulPop(10, allFitnessEffects(genotFitness = rl),</pre>
                         keepPhylog = TRUE, keepEvery = 1,
                         initSize = 4000,
                         seed = NULL, ## for reproducibility
                         mc.cores = 2) ## adapt to your hardware
## Obtain measures of evolutionary predictability
diversityLOD(LOD(simulrl))
## [1] 1.418
diversityPOM(POM(simulrl))
## [1] 1.418
sampledGenotypes(samplePop(simulrl, typeSample = "whole"))
##
    Subjects by Genes matrix of 10 subjects and 6 genes.
##
##
     Genotype Freq
## 1
            \boldsymbol{A}
## 2
             В
         D, F
## 3
                  1
## 4
            \boldsymbol{E}
                  4
## 5
             F
                  3
##
##
   Shannon's diversity (entropy) of sampled genotypes: 1.418
```

#### 1.3.4 Mutator and antimutator genes

The effects of mutator and antimutator genes have been examined both in cancer genetics (Nowak 2006; Tomlinson, Novelli, and Bodmer 1996) and in evolutionary genetics (Gerrish et al. 2007), and are related to wider issues such as Muller's ratchet and the evolution of sex. There are, thus, a large range of questions related to mutator and antimutator genes.

One question addressed in Tomlinson, Novelli, and Bodmer (1996) concerns under what circumstances mutator genes are likely to play a role in cancer progression. For instance, Tomlinson, Novelli, and Bodmer (1996) find that an increased mutation rate is more likely to matter if the number of required mutations in driver genes needed to reach cancer is large and if the mutator effect is large.

We might want to ask, then, how long it takes before to reach cancer under different scenarios. Time to reach cancer is stored in the component FinalTime of the output. We would specify different numbers and effects of mutator genes (argument muEF). We would also change the criteria for reaching cancer and in our case we can easily do that by specifying different numbers in detectionDrivers. Of course, we would also want to examine the effects of varying numbers of mutators, drivers, and possibly fitness consequences of mutators. Below we assume mutators are neutral and we assume there are no additional genes with deleterious mutations, but this need not be so, of course (see also Tomlinson, Novelli, and Bodmer 1996; Gerrish et al. 2007;

Christopher McFarland, Mirny, and Korolev 2014).

Let us run an example. For the sake of simplicity, we assume no epistatic interactions.

Now, simulate using the fitness and mutator specification. We fix the number of drivers to cancer, and we stop when those numbers of drivers are reached. Since we only care about the time it takes to reach cancer, not the actual trajectories, we set keepEvery = NA:

```
## For reproducibility
set.seed(2)
RNGkind("L'Ecuyer-CMRG")
ddr <- 4
st <- oncoSimulPop(4, ft, muEF = mt,
                   detectionDrivers = ddr,
                   finalTime = NA,
                   detectionSize = NA,
                   detectionProb = NA,
                   onlyCancer = TRUE,
                   keepEvery = NA,
                   mc.cores = 2, ## adapt to your hardware
                   seed = NULL) ## for reproducibility
## How long did it take to reach cancer?
unlist(lapply(st, function(x) x$FinalTime))
## [1] 370 141 1793 282
```

(Incidentally, notice that it is easy to get OncoSimulR to throw an exception if you accidentally specify a huge mutation rate when all mutator genes are mutated: see section 14.7.)

# 1.3.5 Epistatic interactions between drivers and passengers in cancer and the consequences of order effects

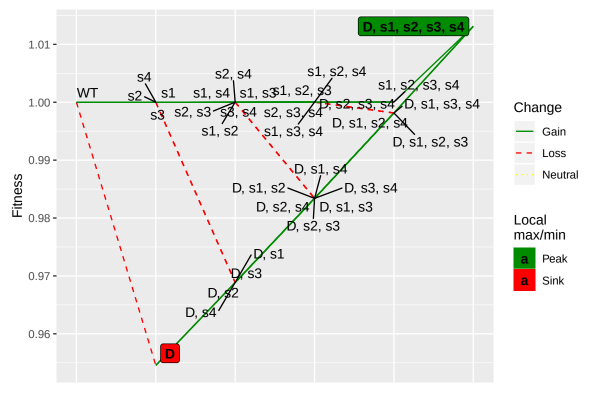
#### 1.3.5.1 Epistatic interactions between drivers and passengers

Bauer, Siebert, and Traulsen (2014) have examined the effects of epistatic relationships between drivers and passengers in cancer initiation. We could use their model as a starting point, and examine how likely cancer is to develop under different variations of their model and different evolutionary scenarios (e.g., initial sample size, mutation rates, evolutionary model, etc).

There are several ways to specify their model, as we discuss in section 5.1. We will use one based on DAGs here:

```
K <- 4
sp <- 1e-5
sdp <- 0.015
sdplus <- 0.05
sdminus <- 0.1
cnt \langle -(1 + sdplus)/(1 + sdminus)
prod_cnt <- cnt - 1</pre>
bauer <- data.frame(parent = c("Root", rep("D", K)),</pre>
                     child = c("D", paste0("s", 1:K)),
                     s = c(prod cnt, rep(sdp, K)),
                     sh = c(0, rep(sp, K)),
                    typeDep = "MN")
fbauer <- allFitnessEffects(bauer)</pre>
(b1 <- evalAllGenotypes(fbauer, order = FALSE, addwt = TRUE))
##
                Genotype Fitness
## 1
                      WT 1.0000
## 2
                       D
                          0.9545
## 3
                      s1
                          1.0000
## 4
                      s2
                          1.0000
## 5
                      s3 1.0000
                         1.0000
## 6
                      s4
## 7
                   D, s1
                         0.9689
## 8
                   D, s2
                         0.9689
## 9
                   D, s3
                          0.9689
## 10
                   D, s4
                         0.9689
## 11
                  s1, s2
                          1.0000
## 12
                  s1, s3 1.0000
## 13
                  s1, s4
                         1.0000
## 14
                  s2, s3
                          1.0000
## 15
                  s2, s4 1.0000
## 16
                  s3, s4 1.0000
## 17
              D, s1, s2 0.9834
              D, s1, s3 0.9834
## 18
```

```
## 19
              D, s1, s4
                          0.9834
              D, s2, s3
## 20
                          0.9834
## 21
              D, s2, s4
                          0.9834
## 22
              D, s3, s4
                          0.9834
## 23
             s1, s2, s3
                          1.0000
## 24
             s1, s2, s4
                          1.0000
## 25
             s1, s3, s4
                          1.0000
## 26
             s2, s3, s4
                          1.0000
          D, s1, s2, s3
## 27
                          0.9981
## 28
          D, s1, s2, s4
                          0.9981
## 29
          D, s1, s3, s4
                          0.9981
## 30
          D, s2, s3, s4
                          0.9981
## 31
         s1, s2, s3, s4
                          1.0000
## 32 D, s1, s2, s3, s4
                          1.0131
## How does the fitness landscape look like?
plot(b1, use_ggrepel = TRUE) ## avoid overlapping labels
```



Now run simulations and examine how frequently the runs end up with population sizes larger than a pre-specified threshold; for instance, below we look at increasing population size 4x in the default maximum number of 2281 time periods (for real, you would of course increase the number of total populations, the range of initial population sizes, model, mutation rate, required population size or number of drivers, etc):

Alternatively, to examine how long it takes to reach cancer for a pre-specified size, you could look at the value of FinalTime as we did above (section 1.3.4) after running simulations with onlyCancer = TRUE and detectionSize set to some reasonable value:

#### 1.3.5.2 Consequences of order effects for cancer initiation

Instead of focusing on different models for epistatic interactions, you might want to examine the consequences of order effects (Ortmann et al. 2015). You would proceed as above, but using models that differ by, say, the presence or absence of order effects. Details on their specification are provided in section 3.6. Here is one particular model (you would, of course, want to compare this to models without order effects or with other magnitudes and types of order effects):

```
## Order effects involving three genes.

## Genotype "D, M" has different fitness effects
## depending on whether M or D mutated first.
```

```
## Ditto for genotype "F, D, M".
## Meaning of specification: X > Y means
## that X is mutated before Y.
o3 <- allFitnessEffects(orderEffects = c(
                             "F > D > M" = -0.3,
                             "D > F > M" = 0.4,
                             "D > M > F" = 0.2,
                             "D > M"
                                         = 0.1,
                             "M > D"
                                         = 0.5))
## With the above specification, let's double check
## the fitness of the possible genotypes
(oeag <- evalAllGenotypes(o3, addwt = TRUE, order = TRUE))</pre>
##
       Genotype Fitness
## 1
             WT
                    1.00
## 2
              D
                    1.00
              F
## 3
                    1.00
## 4
              Μ
                    1.00
## 5
          D > F
                    1.00
## 6
          D > M
                    1.10
          F > D
## 7
                    1.00
## 8
          F > M
                    1.00
## 9
          M > D
                    1.50
## 10
          M > F
                    1.00
## 11 D > F > M
                   1.54
## 12 D > M > F
                    1.32
## 13 F > D > M
                    0.77
## 14 F > M > D
                    1.50
## 15 M > D > F
                    1.50
## 16 M > F > D
                    1.50
```

Now, run simulations and examine how frequently the runs do not end up in extinction. As above, for real, you would of course increase the number of total populations, the range of initial population sizes, mutation rate, etc:

```
mc.cores = 2, ## adapt to your hardware
seed = NULL) ## for reproducibility

## What proportion of the simulations do not end up extinct?
sum(summary(soe1)[, "TotalPopSize"] > 0)/totalpops
## [1] 0.4
```

As we just said, alternatively, to examine how long it takes to reach cancer you could run simulations with onlyCancer = TRUE and look at the value of FinalTime as we did above (section 1.3.4).

#### 1.3.6 Simulating evolution with frequency-dependent fitness

The new frequency-dependent fitness funcionality allows users to run simulations in a different way, defining fitness (birth rates) as functions of clone's frequencies. We can thus model frequency-dependent selection, as well as predation and parasitism, cooperation and mutualism, and commensalism. See section 10 for further details and examples.

#### 1.4 Trade-offs and what is OncoSimulR not well suited for

OncoSimulR is designed for complex fitness specifications and selection scenarios and uses forward-time simulations; the types of questions where OncoSimulR can be of help are discussed in sections 1.2 and 1.3 and running time and space consumption of OncoSimulR are addressed in section 2. You should be aware that **coalescent simulations**, sometimes also called backward-time simulations, are much more efficient for simulating neutral data as well as some special selection scenarios (Yuan et al. 2012; Carvajal-Rodriguez 2010; Hoban, Bertorelle, and Gaggiotti 2011).

In addition, since OncoSimulR allows you to specify fitness with arbitrary epistatic and order effects, as well as mutator effects, you need to learn the syntax of how to specify those effects and you might be paying a performance penalty if your scenario does not require this complexity. For instance, in the model of Beerenwinkel et al. (2007), the fitness of a genotype depends only on the total number of drivers mutated, but not on which drivers are mutated (and, thus, not on the epistatic interactions nor the order of accumulation of the drivers). This means that the syntax for specifying that model could probably be a lot simpler (e.g., specify s per driver).

But it also means that code written for just that case could probably run much faster. First, because fitness evaluation is easier. Second, and possibly much more important, because what we need to keep track of leads to much simpler and economic structures: we do not need to keep track of clones (where two cells are regarded as different clones if they differ anywhere in their genotype), but only of clone types or clone classes as defined by the number of mutated drivers, and keeping track of clones can be expensive —see sections 2 and 14.2.

So for those cases where you do not need the full flexibility of OncoSimulR, special purpose software might be easier to use and faster to run. Of course, for some types

of problems this special purpose software might not be available, though.

### 1.5 Steps for using OncoSimulR

Using this package will often involve the following steps:

- 1. Specify fitness effects: sections 3 and 5.
- 2. Simulate cancer progression: section 6. You can simulate for a single individual or subject or for a set of subjects. You will need to:
  - Decide on a model. This basically amounts to choosing a model with exponential growth ("Exp" or "Bozic") or a model with carrying capacity ("McFL"). If exponential growth, you can choose whether the the effects of mutations operate on the death rate ("Bozic") or the birth rate ("Exp")<sup>1</sup>.
  - Specify other parameters of the simulation. In particular, decide when to stop the simulation, mutation rates, etc.

Of course, at least for initial playing around, you can use the defaults.

3. Sample from the simulated data and do something with those simulated data (e.g., fit an OT model to them, examine diversity or time until cancer, etc). Most of what you do with the data, however, is outside the scope of this package and this vignette.

Before anything else, let us load the package in case it was not yet loaded. We also explicitly load *graph* and *igraph* for the vignette to work (you do not need that for your usual interactive work). And I set the default color for vertices in igraph.

```
library(OncoSimulR)
library(graph)
library(igraph)
igraph_options(vertex.color = "SkyBlue2")
```

To be explicit, what version are we running?

```
packageVersion("OncoSimulR")
## [1] '2.17.996'
```

## 1.6 Two quick examples of fitness specifications

Following 1.5 we will run two very minimal examples. First a model with a few genes and **epistasis**:

```
## 1. Fitness effects: here we specify an
## epistatic model with modules.
```

<sup>&</sup>lt;sup>1</sup>It is of course possible to do this with the carrying capacity (or gompertz-like) models, but there probably is little reason to do it. McFarland et al. (2013) discuss this has little effect on their results, for example. In addition, decreasing the death rate will more easily lead to numerical problems as shown in section 3.11.2.

```
sa <- 0.1
sb < -0.2
sab <- 0.25
sac < -0.1
sbc < -0.25
sv2 <- allFitnessEffects(epistasis = c("-A : B" = sb,</pre>
                                       "A : -B" = sa,
                                       "A : C" = sac,
                                       "A:B" = sab,
                                       "-A:B:C" = sbc),
                         geneToModule = c(
                             "A" = "a1, a2",
                             "B" = "b"
                             "C" = "c"),
                         drvNames = c("a1", "a2", "b", "c"))
evalAllGenotypes(sv2, addwt = TRUE)
##
          Genotype Fitness
## 1
                WT
                     1.000
## 2
                     1.100
                a1
## 3
                   1.100
                a2
                 \boldsymbol{b}
                     0.800
## 4
## 5
                 c 1.000
## 6
           a1, a2
                    1.100
## 7
            a1, b 1.250
## 8
            a1, c 0.990
## 9
             a2, b 1.250
## 10
             a2, c 0.990
## 11
             b, c 1.000
## 12
        a1, a2, b 1.250
## 13
        a1, a2, c 0.990
## 14
          a1, b, c 1.125
## 15
          a2, b, c 1.125
## 16 a1, a2, b, c
                     1.125
## 2. Simulate the data. Here we use the "McFL" model and set
      explicitly parameters for mutation rate, initial size, size
##
      of the population that will end the simulations, etc
RNGkind("Mersenne-Twister")
set.seed(983)
ep1 <- oncoSimulIndiv(sv2, model = "McFL",
                      mu = 5e-6,
                      sampleEvery = 0.025,
                      keepEvery = 0.5,
                      initSize = 2000,
                      finalTime = 3000,
```

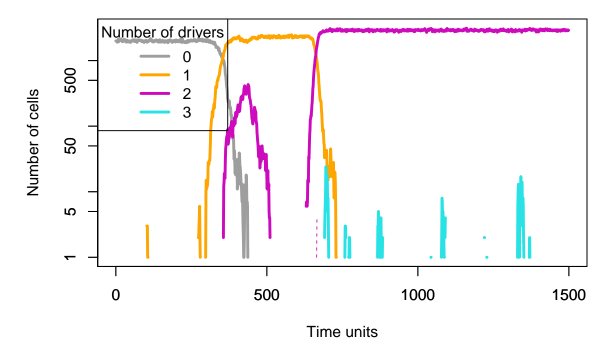


Figure 2: Plot of drivers of an epistasis simulation.

As a second example, we will use a model where we specify **restrictions in the** order of accumulation of mutations using a DAG with the pancreatic cancer poset in Gerstung et al. (2011) (see more details in section 5.5):

```
## Plot the DAG of the fitnessEffects object
plot(pancr)
```

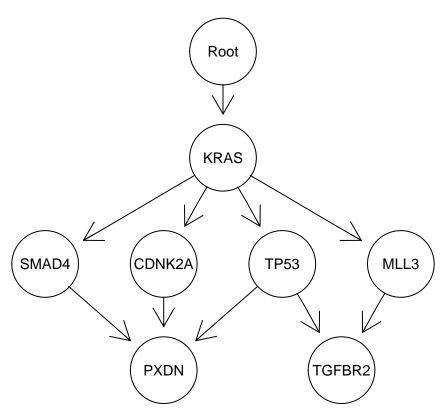


Figure 3: Plot of DAG corresponding to fitnessEffects object.

```
## 2. Simulate from it. We change several possible options.
set.seed(4) ## Fix the seed, so we can repeat it
ep2 <- oncoSimulIndiv(pancr, model = "McFL",
                     mu = 1e-6,
                     sampleEvery = 0.02,
                     keepEvery = 1,
                     initSize = 1000,
                     finalTime = 10000,
                     onlyCancer = FALSE)
## 3. What genotypes and drivers we get? And play with limits
      to show only parts of the data. We also aggressively thin
      the data.
##
par(cex = 0.7)
plot(ep2, show = "genotypes", xlim = c(1000, 8000),
     ylim = c(0, 2400),
     thinData = TRUE, thinData.keep = 0.03)
```

The rest of this vignette explores all of those functions and arguments in much more detail.

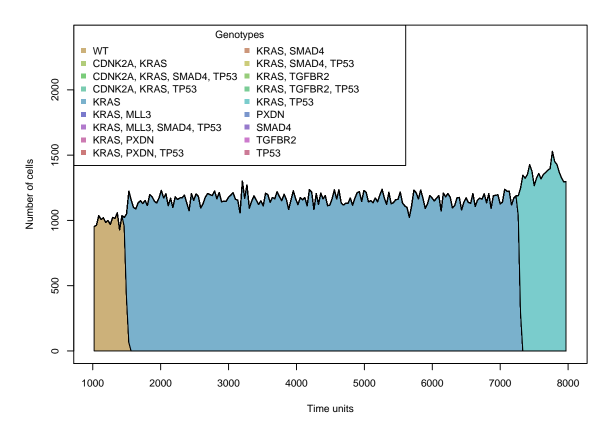


Figure 4: Plot of genotypes of a simulation from a DAG.

# 1.7 Citing OncoSimulR and other documentation

In R, you can do

```
citation("OncoSimulR")
##
## If you use OncoSimulR, please cite the OncoSimulR
## Bioinformatics paper. OncoSimulR has been used in three
## large comparative studies of methods to infer restrictions
## in the order of accumulation of mutations (cancer
## progression models) published in Bioinformatics and BMC
## Bioinformatics; you might want to cite those too, if
## appropriate, such as when referring to using evolutionary
## simulations to assess oncogenetic tree/cancer progression
## methods performance.
##
    R Diaz-Uriarte. OncoSimulR: genetic simulation with
##
##
    arbitrary epistasis and mutator genes in asexual
##
    populations. 2017. Bioinformatics, 33, 1898--1899.
##
    https://doi.org/10.1093/bioinformatics/btx077.
##
    R Diaz-Uriarte. Cancer progression models and fitness
##
     landscapes: a many-to-many relationship 2017
##
```

```
##
                   Bioinformatics.
                    https://academic.oup.com/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/bioinformatics/advance-article/doi/10.1093/
##
##
                   R Diaz-Uriarte. Identifying restrictions in the order of
##
##
                    accumulation of mutations during tumor progression:
                    effects of passengers, evolutionary models, and sampling
##
##
                    2015. BMC Bioinformatics, 16(41).
##
                   R Diaz-Uriarte and C. Vasallo. Every which way? On
##
##
                   predicting tumor evolution using cancer progression models
                   2018 bioRxiv
##
##
                   https://www.biorxiv.org/content/early/2018/11/20/371039
##
## To see these entries in BibTeX format, use
## 'print(<citation>, bibtex=TRUE)', 'toBibtex(.)', or set
## 'options(citation.bibtex.max=999)'.
```

which will tell you how to cite the package. Please, do cite the Bionformatics paper if you use the package in publications.

This is the URL for the Bioinformatics paper: https://doi.org/10.1093/bioinformatics/btx077 (there is also an early preprint at bioRxiv, but it should now point to the Bioinformatics paper).

#### 1.7.1 HTML and PDF versions of the vignette

A PDF version of this vignette is available from https://rdiaz02.github.io/OncoSimul/pdfs/OncoSimulR.pdf. And an HTML version from https://rdiaz02.github.io/OncoSimul/OncoSimulR.html. These files should correspond to the most recent, GitHub version, of the package (i.e., they might include changes not yet available from the BioConductor package).

# 1.8 Testing, code coverage, and other examples

OncoSimulR includes more than 2000 tests that are run at every check cycle. These tests provide a code coverage of more than 90% including both the C++ and R code. Another set of over 500 long-running (several hours) tests can be run on demand (see directory '/tests/manual'). In addition to serving as test cases, some of that code also provides further examples of usage.

#### 1.9 Versions

In this vignette and the documentation I often refer to version 1 (v.1) and version 2 of OncoSimulR. Version 1 is the version available up to, and including, BioConductor v. 3.1. Version 2 of OncoSimulR is available starting from BioConductor 3.2 (and, of course, available too from development versions of BioC). So, if you are using the current stable or development version of BioConductor, or you grab the sources from

GitHub (https://github.com/rdiaz02/OncoSimul) you are using what we call version 2. Please note that the functionality of version 1 will soon be removed.

# 2 Running time and space consumption of OncoSimulR

Time to complete the simulations and size of returned objects (space consumption) depend on several, interacting factors. The usual rule of "experiment before launching a large number of simulations" applies, but here we will walk through several cases to get a feeling for the major factors that affect speed and size. Many of the comments on this section need to use ideas discussed in other places of this document; if you read this section first, you might want to come back after reading the relevant parts.

#### Speed will depend on:

- Your hardware, of course.
- The evolutionary model.
- The granularity of how often you keep data (keepEvery argument). Note that the default, which is to keep as often as you sample (so that we preserve all history) can lead to slow execution times.
- The mutation rate, because higher mutation rates lead to more clones, and more clones means we need to iterate over, well, more clones, and keep larger data structures.
- The fitness specification: more complex fitness specifications tend to be slightly slower but specially different fitness specifications can have radically different effects on the evolutionary trajectories, accessibility of fast growing genotypes and, generally, the evolutionary dynamics.
- The stopping conditions (detectionProb, detectionDrivers, detectionSize arguments) and whether or not simulations are run until cancer is reached (onlyCancer argument).
- Most of the above factors can interact in complex ways.

#### Size of returned objects will depend on:

- Any factor that affects the number of clones tracked/returned, in particular: initial sizes and stopping conditions, mutation rate, and how often you keep data (the keepEvery argument can make a huge difference here).
- Whether or not you keep the complete genealogy of clones (this affects slightly the size of returned object, not speed).

In the sections that follow, we go over several cases to understand some of the main settings that affect running time (or execution time) and space consumption (the size of returned objects). It should be understood, however, that many of the examples shown below do not represent typical use cases of OncoSimulR and are used only to identify what and how affects running time and space consumption. As we will see in most examples in this vignette, typical use cases of OncoSimulR involve hundreds to thousands of genes on population sizes up to  $10^5$  to  $10^7$ .

Note that most of the code in this section is not executed during the building of the vignette to keep vignette build time reasonable and prevent using huge amounts of RAM. All of the code, ready to be sourced and run, is available from the 'inst/miscell' directory (and the summary output from some of the benchmarks is available from

the 'miscell-files/vignette\_bench\_Rout' directory of the main OncoSimul repository at https://github.com/rdiaz02/OncoSimul).

# 2.1 Exp and McFL with "detectionProb" and pancreas example

To get familiar with some of they factors that affect time and size, we will use the fitness specification from section 1.6, with the detectionProb stopping mechanism (see 6.3.2). We will use the two main growth models (exponential and McFarland). Each model will be run with two settings of keepEvery. With keepEvery = 1 (runs exp1 and mc1), population samples are stored at time intervals of 1 (even if most of the clones in those samples later become extinct). With keepEvery = NA (runs exp2 and mc2) no intermediate population samples are stored, so clones that become extinct at any sampling period are pruned and only the existing clones at the end of the simulation are returned (see details in 14.2.1).

Will run 100 simulations. The results I show are for a laptop with an 8-core Intel Xeon E3-1505M CPU, running Debian GNU/Linux (the results from these benchmarks are available as data(benchmark 1)).

```
## Specify fitness
pancr <- allFitnessEffects(</pre>
    data.frame(parent = c("Root", rep("KRAS", 4),
                    "SMAD4", "CDNK2A",
                    "TP53", "TP53", "MLL3"),
               child = c("KRAS", "SMAD4", "CDNK2A",
                    "TP53", "MLL3",
                    rep("PXDN", 3), rep("TGFBR2", 2)),
               s = 0.1,
               sh = -0.9,
               typeDep = "MN"),
    drvNames = c("KRAS", "SMAD4", "CDNK2A", "TP53",
                  "MLL3", "TGFBR2", "PXDN"))
Nindiv <- 100 ## Number of simulations run.
              ## Increase this number to decrease sampling variation
## keepEvery = 1
t_exp1 <- system.time(</pre>
    exp1 <- oncoSimulPop(Nindiv, pancr,</pre>
                             detectionProb = "default",
                             detectionSize = NA,
                             detectionDrivers = NA,
                             finalTime = NA,
                             keepEvery = 1,
                             model = "Exp",
                             mc.cores = 1))["elapsed"]/Nindiv
```

```
t mc1 <- system.time(</pre>
    mc1 <- oncoSimulPop(Nindiv, pancr,</pre>
                             detectionProb = "default",
                             detectionSize = NA,
                             detectionDrivers = NA,
                             finalTime = NA,
                             keepEvery = 1,
                             model = "McFL",
                             mc.cores = 1))["elapsed"]/Nindiv
## keepEvery = NA
t_exp2 <- system.time(</pre>
    exp2 <- oncoSimulPop(Nindiv, pancr,</pre>
                              detectionProb = "default",
                              detectionSize = NA,
                              detectionDrivers = NA,
                              finalTime = NA,
                              keepEvery = NA,
                              model = "Exp",
                              mc.cores = 1))["elapsed"]/Nindiv
t mc2 <- system.time(</pre>
    mc2 <- oncoSimulPop(Nindiv, pancr,</pre>
                             detectionProb = "default",
                             detectionSize = NA,
                             detectionDrivers = NA,
                             finalTime = NA,
                             keepEvery = NA,
                             model = "McFL",
                             mc.cores = 1))["elapsed"]/Nindiv
```

We can obtain times, sizes of objects, and summaries of numbers of clones, iterations, and final times doing, for instance:

```
cat("\n\n\n t_exp1 = ", t_exp1, "\n")
object.size(exp1)/(Nindiv * 1024^2)
cat("\n\n")
summary(unlist(lapply(exp1, "[[", "NumClones")))
summary(unlist(lapply(exp1, "[[", "NumIter")))
summary(unlist(lapply(exp1, "[[", "FinalTime")))
summary(unlist(lapply(exp1, "[[", "TotalPopSize")))
```

The above runs yield the following:

Table 2: Benchmarks of Exp and McFL models using the default detectionProb with two settings of keepEvery.

	Elapsed Time, average per simulation (s)	Object Size, average per simulation (MB)	Number of Clones, median	Number of Iterations, median	Final Time, median	Total Population Size, median	Total Population Size, max.	keepEvery
exp1	0	0.04	2	254	252	1,058	11,046	1
mc1	0.74	3.9	12	816,331	20,406	696	979	1
$\mathbf{exp2}$	0	0.01	1	296	294	1,021	21,884	NA
mc2	0.7	0.01	1	694,716	$17,\!366$	692	888	NA

The above table shows that a naive comparison (looking simply at execution time) might conclude that the McFL model is much, much slower than the Exp model. But that is not the complete story: using the detectionProb stopping mechanism (see 6.3.2) will lead to stopping the simulations very quickly in the exponential model because as soon as a clone with fitness > 1 appears it starts growing exponentially. In fact, we can see that the number of iterations and the final time are much smaller in the Exp than in the McFL model. We will elaborate on this point below (section 2.2.1), when we discuss the setting for checkSizePEvery (here left at its default value of 20): checking the exiting condition more often (smaller checkSizePEvery) would probably be justified here (notice also the very large final times) and would lead to a sharp decrease in number of iterations and, thus, running time.

This table also shows that the keepEvery = NA setting, which was in effect in simulations exp2 and mc2, can make a difference especially for the McFL models, as seen by the median number of clones and the size of the returned object. Models exp2 and mc2 do not store any intermediate population samples so clones that become extinct at any sampling period are pruned and only the existing clones at the end of the simulation are returned. In contrast, models exp1 and mc1 store population samples at time intervals of 1 (keepEvery = 1), even if many of those clones eventually become extinct. We will return to this issue below as execution time and object size depend strongly on the number of clones tracked.

We can run the exponential model again modifying the arguments of the detectionProb mechanism; in two of the models below (exp3 and exp4) no detection can take place unless populations are at least 100 times larger than the initial population size, and probability of detection is 0.1 with a population size 1,000 times larger than the initial one (PDBaseline = 5e4, n2 = <math>5e5). In the other two models (exp5 and exp6), no detection can take place unless populations are at least 1,000 times larger than the initial population size, and probability of detection is 0.1 with a population size 100,000 times larger than the initial one (PDBaseline = 5e5, n2 = <math>5e7)<sup>2</sup>. In runs exp3 and exp5 we set keepEvery = 1 and in runs exp4 and exp6 we set keepEvery = NA.

 $<sup>^2</sup>$ Again, these are not necessarily reasonable or common settings. We are using them to understand what and how affects running time and space consumption.

```
t_exp4 <- system.time(</pre>
    exp4 <- oncoSimulPop(Nindiv, pancr,</pre>
                              detectionProb = c(PDBaseline = 5e4,
                                                 p2 = 0.1, n2 = 5e5,
                                                 checkSizePEvery = 20),
                              detectionSize = NA,
                              detectionDrivers = NA,
                             finalTime = NA,
                             keepEvery = NA,
                             model = "Exp",
                             mc.cores = 1))["elapsed"]/Nindiv
t exp5 <- system.time(</pre>
    exp5 <- oncoSimulPop(Nindiv, pancr,</pre>
                              detectionProb = c(PDBaseline = 5e5,
                                                p2 = 0.1, n2 = 5e7),
                             detectionSize = NA,
                              detectionDrivers = NA,
                             finalTime = NA,
                             keepEvery = 1,
                             model = "Exp",
                             mc.cores = 1))["elapsed"]/Nindiv
t_exp6 <- system.time(</pre>
    exp6 <- oncoSimulPop(Nindiv, pancr,</pre>
                              detectionProb = c(PDBaseline = 5e5,
                                                p2 = 0.1, n2 = 5e7),
                              detectionSize = NA,
                              detectionDrivers = NA,
                             finalTime = NA,
                             keepEvery = NA,
                             model = "Exp",
                             mc.cores = 1))["elapsed"]/Nindiv
```

Table 3: Benchmarks of Exp and McFL models modifying the default detectionProb with two settings of keepEvery.

	Elapsed Time, average per simulation (s)	Object Size, average per simulation (MB)	Number of Clones, median	Number of Iterations, median	Final Time, median	Total Population Size, median	Total Population Size, max.	keepE	vePDBaselin2e
exp3	0.01	0.41	14	2,754	1,890	6,798,358	2.7e + 08	1	50,000  5e + 05
$\exp 4$	0.01	0.02	8	2,730	2,090	7,443,812	1.7e + 08	NA	50,000  5e+05
$\exp 5$	0.84	0.91	34	54,332	2,026	1.4e+09	4.2e+10	1	5e+05 5e+07
exp6	0.54	0.02	27	44,288	2,026	1.2e + 09	3.3e + 10	NA	5e+05 $5e+07$

As above, keepEvery = NA (in exp4 and exp6) leads to much smaller object sizes and slightly smaller numbers of clones and execution times. Changing the exiting conditions (by changing detectionProb arguments) leads to large increases in number of iterations (in this case by factors of about 15x to 25x) and a corresponding increase in execution time as well as much larger population sizes (in some cases  $> 10^{10}$ ).

In some of the runs of exp5 and exp6 we get the (recoverable) exception message from the C++ code: Recoverable exception ti set to DBL\_MIN. Rerunning, which is related to those simulations reaching total population sizes  $> 10^{10}$ ; we return to this below (section 2.4). You might also wonder why total and median population sizes are so large in these two runs, given the exiting conditions. One of the reasons is that we are using the default checkSizePEvery = 20, so the interval between successive checks of the exiting condition is large; this is discussed at greater length in section 2.2.1.

All the runs above used the default value onlyCancer = TRUE. This means that simulations will be repeated until the exiting conditions are reached (see details in section 6.3) and, therefore, any simulation that ends up in extinction will be repeated. This setting can thus have a large effect on the exponential models, because when the initial population size is not very large and we start from the wildtype, it is not uncommon for simulations to become extinct (when birth and death rates are equal and the population size is small, it is easy to reach extinction before a mutation in a gene that increases fitness occurs). But this is rarely the case in the McFarland model (unless we use really tiny initial population sizes) because of the dependency of death rate on total population size (see section 3.2.1).

The number of attempts until cancer was reached in the above models is shown in Table 4 (the values can be obtained from any of the above runs doing, for instance, median(unlist(lapply(exp1, function(x) x\$other\$attemptsUsed)))):

-					
	Attempts until	Attempts until	Attempts until		
	Cancer, median	Cancer, mean	Cancer, max.	PDBasel	ine n2
$\overline{\exp 1}$	1	1.9	7	600	1,000
mc1	1	1	1	600	1,000
$\exp 2$	2	2.2	16	600	1,000
mc2	1	1	1	600	1,000
$\exp 3$	6	7.7	40	50,000	5e + 05
exp4	6	8	39	50,000	5e + 05
exp5	5	8.3	41	5e + 05	5e + 07
$\exp 6$	5	7.2	30	5e + 05	5e + 07

Table 4: Number of attempts until cancer.

The McFL models finish in a single attempt. The exponential model simulations where we can exit with small population sizes (exp1, exp2) need many fewer attempts to reach cancer than those where large population sizes are required (exp3 to exp6).

There is no relevant different among those last four, which is what we would expect: a population that has already reached a size of 50,000 cells from an initial population size of 500 is obviously a growing population where there is at least one mutant with positive fitness; thus, it unlikely to go extinct and therefore having to grow up to at least 500,000 will not significantly increase the risk of extinction.

We will now rerun all of the above models with argument onlyCancer = FALSE. The results are shown in Table 5 (note that the differences between this table and Table 2 for the McFL models are due only to sampling variation).

Table 5: Benchmarks of models in Table 2 and 3 when run with onlyCancer = FALSE.

	Elapsed										
	Time,	Object Size,				Total	Total	Total			
	average per	average per	Number of	Number of		Population	Popula-	Popula-			
	simulation	simulation	Clones,	Iterations,	Final Time,	Size,	tion Size,	tion Size,			
	(s)	(MB)	median	median	median	median	mean	max.	keepEvery	PDBaseline	n2
exp1_noc	0.001	0.041	1.5	394	393	0	708	18,188	1	600	1,000
$mc1\_noc$	0.69	3.9	12	673,910	16,846	692	700	983	1	600	1,000
$\exp 2\_{noc}$	0.001	0.012	1	320	319	726	870	26,023	NA	600	1,000
$mc2\_noc$	0.65	0.014	1	628,683	15,716	694	704	910	NA	600	1,000
$\exp 3\_noc$	0.002	0.15	2	718	694	0	$2,\!229,\!519$	5.7e + 07	1	50,000	5e + 05
$\exp 4\_noc$	0.002	0.013	0	600	599	0	$3,\!122,\!765$	1.3e + 08	NA	50,000	5e + 05
$\exp 5\_{noc}$	0.17	0.22	3	848	777	0	5.9e + 08	$1.5e{+10}$	1	5e + 05	5e + 07
$\exp6\_noc$	0.068	0.013	0	784	716	0	4.1e+08	$1.3e{+}10$	NA	5e + 05	5e + 07

Now most simulations under the exponential model end up in extinction, as seen by the median population size of 0 (but not all, as the mean and max. population size are clearly away from zero). Consequently, simulations under the exponential model are now faster (and the size of the average returned object is smaller). Of course, whether one should run simulations with onlyCancer = TRUE or onlyCancer = FALSE will depend on the question being asked (see, for example, section 1.3.5 for a question where we will naturally want to use onlyCancer = FALSE).

To make it easier to compare results with those of the next section, Table 6 shows all the runs so far.

Table 6: Benchmarks of all models in Tables 2, 3, and 5.

	Elapsed Time,											
	average	Object Size,				Total						
	per simu-	average per	Number of	Number of		Population	Total	Total				
	lation	simulation	Clones,	Iterations,	Final Time,	Size,	Population	Population				
	(s)	(MB)	median	median	median	median	Size, mean	Size, max.	keepEver	y PDBaseli	ine n2	onlyCancer
exp1	0.001	0.037	2	254	252	1,058	1,277	11,046	1	600	1,000	TRUE
mc1	0.74	3.9	12	816,331	20,406	696	702	979	1	600	1,000	TRUE
$\mathbf{exp2}$	0.001	0.012	1	296	294	1,021	1,392	21,884	NA	600	1,000	TRUE
mc2	0.7	0.014	1	694,716	17,366	692	698	888	NA	600	1,000	TRUE
exp3	0.01	0.41	14	2,754	1,890	6,798,358	1.7e + 07	2.7e + 08	1	50,000	5e + 05	TRUE
$\exp 4$	0.009	0.016	8	2,730	2,090	7,443,812	1.5e + 07	1.7e + 08	NA	50,000	5e + 05	TRUE
exp5	0.84	0.91	34	54,332	2,026	1.4e + 09	3.5e + 09	$4.2e{+}10$	1	5e + 05	5e + 07	TRUE
exp6	0.54	0.021	27	44,288	2,026	1.2e + 09	3.2e + 09	$3.3e{+}10$	NA	5e + 05	5e + 07	TRUE
exp1_noc	0.001	0.041	1.5	394	393	0	708	18,188	1	600	1,000	FALSE
mc1_noc	0.69	3.9	12	673,910	16,846	692	700	983	1	600	1,000	FALSE
$\exp 2$ _noc	0.001	0.012	1	320	319	726	870	26,023	NA	600	1,000	FALSE
$mc2\_noc$	0.65	0.014	1	$628,\!683$	15,716	694	704	910	NA	600	1,000	FALSE
exp3_noc	0.002	0.15	2	718	694	0	$2,\!229,\!519$	5.7e + 07	1	50,000	5e + 05	FALSE
exp4_noc	0.002	0.013	0	600	599	0	$3,\!122,\!765$	1.3e + 08	NA	50,000	5e + 05	FALSE
$\exp 5$ _noc	0.17	0.22	3	848	777	0	5.9e + 08	1.5e + 10	1	5e + 05	5e + 07	FALSE
exp6_noc	0.068	0.013	0	784	716	0	4.1e + 08	$1.3e{+10}$	NA	5e + 05	5e + 07	FALSE

#### **2.1.1** Changing fitness: s = 0.1 and s = 0.05

In the above fitness specification the fitness effect of each gene (when its restrictions are satisfied) is s=0.1 (see section 3.2 for details). Here we rerun all the above benchmarks using s=0.05 (the results from these benchmarks are available as data(benchmark\_1\_0.05)) and results are shown below in Table 7.

Table 7: Benchmarks of all models in Table 6 using s=0.05 (instead of s=0.1).

	Elapsed Time,											
	average	Object Size,				Total						
	per simu-	average per	Number of	Number of		Population	Total	Total				
	lation	simulation	Clones,	Iterations,	Final Time,	Size,	Population	Population				
	(s)	(MB)	median	median	median	median	Size, mean	Size, max.	keepEver	y PDBaseli	ne n2	onlyCancer
exp1	0.002	0.043	2	316	315	1,104	1,181	3,176	1	600	1,000	TRUE
mc1	1.7	11	17	2e + 06	50,696	644	647	761	1	600	1,000	TRUE
$\mathbf{exp2}$	0.001	0.012	1	274	273	1,129	1,281	7,608	NA	600	1,000	TRUE
mc2	1.6	0.016	1	$1,\!615,\!197$	$40,\!376$	644	651	772	NA	600	1,000	TRUE
exp3	0.012	0.63	15	3,995	2,919	3,798,540	5,892,376	4.5e + 07	1	50,000	5e + 05	TRUE
$\exp 4$	0.011	0.017	9	4,288	$3,\!276$	4,528,072	6,551,319	3.2e + 07	NA	50,000	5e + 05	TRUE
exp5	0.3	1.2	34	68,410	2,751	6.8e + 08	1e+09	8.2e + 09	1	5e + 05	5e + 07	TRUE
$\exp 6$	0.26	0.022	23	44,876	2,499	4.3e + 08	8.9e + 08	7.3e + 09	NA	5e + 05	5e + 07	TRUE
exp1_noc	0.001	0.039	2	310	308	0	522	2,239	1	600	1,000	FALSE
mc1_noc	1.6	11	17	2e + 06	50,776	638	643	757	1	600	1,000	FALSE
$\exp 2$ _noc	0.001	0.012	0	340	336	0	599	3,994	NA	600	1,000	FALSE
$mc2\_noc$	1.7	0.017	1	2,102,439	$52,\!556$	645	650	740	NA	600	1,000	FALSE
exp3_noc	0.002	0.11	2	618	615	0	150,978	6,093,498	1	50,000	5e + 05	FALSE
exp4_noc	0.002	0.013	0	813	812	0	$558,\!225$	2.3e + 07	NA	50,000	5e + 05	FALSE
$\exp 5$ _noc	0.031	0.23	3	917	914	0	1.1e + 08	3.7e + 09	1	5e + 05	5e + 07	FALSE
exp6_noc	0.046	0.013	0	628	610	0	1.7e + 08	5.1e + 09	NA	5e + 05	5e + 07	FALSE

As expected, having a smaller s leads to slower processes in most cases, since it takes longer to reach the exiting conditions sooner. Particularly noticeable are the runs for the McFL models (notice the increases in population size and number of iterations—see also below).

That is not the case, however, for exp5 and exp6 (and exp5\_noc and exp6\_noc). When running with s=0.05 the simulations exit at a later time (see column "Final Time") but they exit with smaller population sizes. Here we have an interaction between sampling frequency, speed of growth of the population, mutation events and number of clones. In populations that grow much faster mutation events will happen more often (which will trigger further iterations of the algorithm); in addition, more new clones will be created, even if they only exist for short times and become extinct by the following sampling period (so they are not reflected in the pops.by.time matrix). These differences are proportionally larger the larger the rate of growth of the population. Thus, they are larger between, say, the exp5 at s=0.1 and s=0.05 than between the exp4 at the two different s: the exp5 exit conditions can only be satisfied at much larger population sizes so at populations sizes when growth is much faster (recall we are dealing with exponential growth).

Recall also that with the default settings in detectionProb, we assess the exiting condition every 20 time periods (argument checkSizePEvery); this means that for fast growing populations, the increase in population size between successive checks of the exit conditions will be much larger (this phenomenon is also discussed in section 2.2.1).

Thus, what is happening in the exp5 and exp6 with s=0.1 is that close to the time the exit conditions could be satisfied, they are growing very fast, accumulating mutants, and incurring in additional iterations. They exit sooner in terms of time periods, but they do much more work before arriving there.

The setting of checkSizePEvery is also having a huge effect on the McFL model simulations (the number of iterations is  $> 10^6$ ). Even more than in the previous section, checking the exiting condition more often (smaller checkSizePEvery) would probably be justified here (notice also the very large final times) and would lead to a sharp decrease in number of iterations and, thus, running time.

The moral here is that in complex simulations like this (and most simulations are complex), the effects of some parameters (s in this case) might look counter-intuitive at first. Thus the need to "experiment before launching a large number of simulations".

#### 2.2 Several "common use cases" runs

Let us now execute some simulations under more usual conditions. We will use seven different fitness specifications: the pancreas example, two random fitness landscapes, and four sets of independent genes (200 to 4000 genes) with fitness effects randomly drawn from exponential distributions:

```
pancr <- allFitnessEffects(</pre>
    data.frame(parent = c("Root", rep("KRAS", 4),
                   "SMAD4", "CDNK2A",
                   "TP53", "TP53", "MLL3"),
               child = c("KRAS", "SMAD4", "CDNK2A",
                   "TP53", "MLL3",
                   rep("PXDN", 3), rep("TGFBR2", 2)),
               s = 0.1,
               sh = -0.9,
               typeDep = "MN"),
    drvNames = c("KRAS", "SMAD4", "CDNK2A", "TP53",
                 "MLL3", "TGFBR2", "PXDN"))
## Random fitness landscape with 6 genes
## At least 50 accessible genotypes
rf16 <- rfitness(6, min_accessible_genotypes = 50)</pre>
attributes(rf16) $accessible genotypes ## How many accessible
rf6 <- allFitnessEffects(genotFitness = rf16)</pre>
## Random fitness landscape with 12 genes
## At least 200 accessible genotypes
rfl12 <- rfitness(12, min_accessible_genotypes = 200)
attributes(rf112) $accessible genotypes ## How many accessible
rf12 <- allFitnessEffects(genotFitness = rf112)</pre>
## Independent genes; positive fitness from exponential distribution
## with mean around 0.1, and negative from exponential with mean
## around -0.02. Half of genes positive fitness effects, half
## negative.
ng <- 200 re 200 <- allFitnessEffects(noIntGenes = c(rexp(ng/2, 10),
-rexp(ng/2, 50))
ng <- 500
re_500 <- allFitnessEffects(noIntGenes = c(rexp(ng/2, 10),
                                            -rexp(ng/2, 50))
ng <- 2000
re 2000 <- allFitnessEffects(noIntGenes = c(rexp(ng/2, 10),
                                             -rexp(ng/2, 50))
```

#### 2.2.1 Common use cases, set 1.

We will use the Exp and the McFL models, run with different parameters. The script is provided as 'benchmark\_2.R', under '/inst/miscell', with output in the 'miscell-files/vignette\_bench\_Rout' directory of the main OncoSimul repository at https://github.com/rdiaz02/OncoSimul. The data are available as data(benchmark\_2).

For the Exp model the call will be

```
oncoSimulPop(Nindiv,
    fitness,
    detectionProb = NA,
    detectionSize = 1e6,
    initSize = 500,
    detectionDrivers = NA,
    keepPhylog = TRUE,
    model = "Exp",
    errorHitWallTime = FALSE,
    errorHitMaxTries = FALSE,
    finalTime = 5000,
    onlyCancer = FALSE,
    mc.cores = 1,
    sampleEvery = 0.5,
    keepEvery = 1)
```

And for McFL:

```
initSize <- 1000
oncoSimulPop(Nindiv,
              fitness,
               detectionProb = c(
                   PDBaseline = 1.4 * initSize,
                   n2 = 2 * initSize,
                   p2 = 0.1,
                   checkSizePEvery = 4),
               initSize = initSize,
               detectionSize = NA,
               detectionDrivers = NA,
               keepPhylog = TRUE,
               model = "McFL",
               errorHitWallTime = FALSE,
               errorHitMaxTries = FALSE,
               finalTime = 5000,
```

```
max.wall.time = 10,
onlyCancer = FALSE,
mc.cores = 1,
keepEvery = 1)
```

For the exponential model we will stop simulations when populations have > 10<sup>6</sup> cells (simulations start from 500 cells). For the McFarland model we will use the detectionProb mechanism (see section 6.3.2 for details); we could have used as stopping mechanism detectionSize = 2 \* initSize (which would be basically equivalent to reaching cancer, as argued in (McFarland et al. 2013)) but we want to provide further examples under the detectionProb mechanism. We will start from 1000 cells, not 500 (starting from 1000 we almost always reach cancer in a single run).

Why not use the detectionProb mechanism with the Exp models? Because it can be hard to intuitively understand what are reasonable settings for the parameters of the detectionProb mechanism when used in a population that is growing exponentially, especially if different genes have very different effects on fitness. Moreover, we are using fitness specifications that are very different (compare the fitness landscape of six genes, the pancreas specification, and the fitness specification with 4000 genes with fitness effects drawn from an exponential distribution —re\_4000). In contrast, the detectionProb mechanism might be simpler to reason about in a population that is growing under a model of carrying capacity with possibly large periods of stasis. Let us emphasize that it is not that the detectionProb mechanism does not make sense with the Exp model; it is simply that the parameters might need finer adjustment for them to make sense, and in these benchmarks we are dealing with widely different fitness specifications.

Note also that we specify <code>checkSizePEvery = 4</code> (instead of the default, which is 20). Why? Because the fitness specifications where fitness effects are drawn from exponential distributions (<code>re\_200</code> to <code>re\_4000</code> above) include many genes (well, up to 4000) some of them with possibly very large effects. In these conditions, simulations can run very fast in the sense of "units of time". If we check exiting conditions every 20 units the population could have increased its size several orders of magnitude in between checks (this is also discussed in sections 2.1.1 and 6.3.2). You can verify this by running the script with other settings for <code>checkSizePEvery</code> (and being aware that large settings might require you to wait for a long time). To ensure that populations have really grown, we have increased the setting of <code>PDBaseline</code> so that no simulation can be considered for stopping unless its size is 1.4 times larger than <code>initSize</code>.

In all cases we use keepEvery = 1 and keepPhylog = TRUE (so we store the population sizes of all clones every 1 time unit and we keep the complete genealogy of clones). Finally, we run all models with errorHitWallTime = FALSE and errorHitMaxTries = FALSE so that we can see results even if stopping conditions are not met.

The results of the benchmarks, using 100 individual simulations, are shown in Table 8.

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Table 8: Benchmarks under some common use cases, set 1.

ModelFitness	Elapsed Time, average per simulation (s)	Object Size, average per simulation (MB)	Number of Clones, median	Number of Iterations, median	Final Time, median	Total Population Size, median	Total Population Size, mean	Total Population Size, max.
Exp pancr	0.002	0.12	3	1,397	697	0	164,222	1,053,299
McFLpancr	0.12	0.56	8	2e + 05	5,000	1,037	1,144	1,938
Exp rf6	0.002	0.064	6	783	391	1e+06	594,899	1,309,497
McFLrf6	0.019	0.071	3	$23,\!297$	582	1,884	1,975	4,636
Exp rf12	0.01	0.13	4	1,178	542	0	287,669	1,059,141
McFLrf12	0.14	0.82	18	2e + 05	5,000	1,252	1,295	1,695
$Exp re_200$	0.013	0.67	230	1,185	223	1,060,944	859,606	1,536,242
$McFLre\_200$	0.018	0.22	47	9,679	240	2,166	2,973	29,301
$Exp re_500$	0.09	2.7	771	2,732	152	1,068,732	959,026	1,285,522
$McFLre\_500$	0.024	0.44	91	7,056	172	2,148	2,578	8,234
$Exp re_2000$	0.91	29	3,376	$7,\!412$	70	1,163,990	1,143,041	1,741,492
$McFLre\_2000$	0.031	1.9	186	3,546	80	2,870	3,704	13,248
$\rm Exp \ re\_4000$	3.3	113	7,088	12,216	52	1,217,568	1,309,185	2,713,200
$McFLre\_4000$	0.063	6.5	326	2,731	52	4,592	13,601	729,611

In most cases, simulations run reasonably fast (under 0.1 seconds per individual simulation) and the returned objects are small. I will only focus on a few cases.

The McFL model with random fitness landscape rf12 and with pancr does not satisfy the conditions of detectionProb in most cases: its median final time is 5000, which was the maximum final time specified. This suggests that the fitness landscape is such that it is unlikely that we will reach population sizes > 1400 (remember we the setting for PDBaseline) before 5000 time units. There is nothing particular about using a fitness landscape of 12 genes and other runs in other 12-gene random fitness landscapes do not show this pattern. However, complex fitness landscapes might be such that genotypes of high fitness (those that allow reaching a large population size quickly) are not easily accessible<sup>3</sup> so reaching them might take a long time. This does not affect the exponential model in the same way because, well, because there is exponential growth in that model: any genotype with fitness > 1will grow exponentially (of course, at possibly very different rates). You might want to play with the script and modify the call to rfitness (using different values of reference and c, for instance) to have simpler paths to a maximum or modify the call to oncoSimulPop (with, say, finalTime to much larger values). Some of these issues are related to more general questions about fitness landscapes and accessibility (see section 1.3.2 and references therein).

You could also set onlyCancer = TRUE. This might make sense if you are interested in only seeing simulations that "reach cancer" (where "reach cancer" means reaching a state you define as a function of population size or drivers). However, if you are exploring fitness landscapes, onlyCancer = TRUE might not always be reasonable as reaching a particular population size, for instance, might just not be possible under some fitness landscapes (this phenomenon is of course not restricted to random fitness landscapes —see also section 2.3.3).

As we anticipated above, the detectionProb mechanism has to be used with care: some of the simulations run in very short "time units", such as those for the fitness specifications with 2000 and 4000 genes. Having used a checkSizePEvery = 20 probably would not have made sense.

Finally, it is interesting that in the cases examined here, the two slowest running simulations are from "Exp", with fitnesses  $re_2000$  and  $re_4000$  (and the third slowest is also Exp, under  $re_500$ ). These are also the cases with the largest number of clones. Why? In the "Exp" model there is no competition, and fitness specifications  $re_2000$  and  $re_4000$  have genomes with many genes with positive fitness contributions. It is thus very easy to obtain, from the wildtype ancestor, a large number of clones all of which have birth rates > 1 and, thus, clones that are unlikely to become extinct.

<sup>&</sup>lt;sup>3</sup>By easily accessible I mean that there are many, preferably short, paths of non-decreasing fitness from the wildtype to this genotype. See definitions and discussion in, e.g., Franke et al. (2011).

#### 2.2.2 Common use cases, set 2.

We will now rerun the simulations above changing the following:

- finalTime set to 25000.
- onlyCancer set to TRUE.
- The "Exp" models will stop when population size  $> 10^5$ .

This is in script 'benchmark\_3.R', under '/inst/miscell', with output in the 'miscell-files/vignette\_bench\_Rout' directory of the main OncoSimul repository at https://github.com/rdiaz02/OncoSimul. The data are available as data(benchmark\_3).

Table 9: Benchmarks under some common use cases, set 2.

ModelFitness	Elapsed Time, average per simulation (s)	Object Size, average per simulation (MB)	Number of Clones, median	Number of Iterations, median	Final Time, median	Total Population Size, median	Total Population Size, mean	Total Population Size, max.
Exp pancr	0.012	0.32	10	3,480	1,718	1e+05	1e+05	108,805
McFLpancr	0.41	1.7	14	4e + 05	9,955	1,561	1,555	1,772
Exp rf6	0.003	0.058	4	866	430	107,492	109,774	$135,\!257$
McFLrf6	0.033	0.12	4	35,216	880	2,003	2,010	3,299
Exp rf12	0.012	0.098	9	1,138	561	1e + 05	1e+05	112,038
McFLrf12	0.17	0.76	16	1e+05	2,511	1,486	1,512	1,732
$Exp re_200$	0.004	0.39	106	723	252	1e + 05	$105,\!586$	122,338
$McFLre\_200$	0.026	0.33	61	13,484	335	1,830	2,049	3,702
$Exp re_500$	0.007	0.61	168	490	117	110,311	$112,\!675$	134,860
$McFLre\_500$	0.018	0.33	70	5,157	126	2,524	3,455	19,899
$Exp re_2000$	0.046	5.7	651	1,078	68	106,340	109,081	153,146
$McFLre\_2000$	0.029	1.8	186	3,444	80	2,837	4,009	37,863
$Exp re\_4000$	0.1	19	1,140	1,722	51	111,256	113,499	168,958
$McFLre\_4000$	0.057	6.7	325	3,081	60	3,955	8,892	$265,\!183$

Since we increased the maximum final time and forced runs to "reach cancer" the McFL run with the pancreas fitness specification takes a bit longer because it also has to do a larger number of iterations. Interestingly, notice that the median final time is close to 10000, so the runs in 2.2.1 with maximum final time of 5000 would have had a hard time finishing with onlyCancer = TRUE.

Forcing simulations to "reach cancer" and just random differences between the random fitness landscape also affect the McFL run under rf12: final time is below 5000 and the median number of iterations is about half of what was above.

Finally, by stopping the Exp simulations at 10<sup>5</sup>, simulations with re\_2000 and re\_4000 finish now in much shorter times (but they still take longer than their McFL counterparts) and the number of clones created is much smaller.

#### 2.3 Can we use a large number of genes?

Yes. In fact, in OncoSimulR there is no pre-set limit on genome size. However, large numbers of genes can lead to unacceptably large returned object sizes and/or running time. We discuss several examples next that illustrate some of the major issues to consider. Another example with 50,000 genes is shown in section 6.5.3.

We have seen in 2.1 and 2.2.1 that for the Exp model, benchmark results using detectionProb require a lot of care and can be misleading. Here, we will fix initial population sizes (to 500) and all final population sizes will be set to  $\geq 10^6$ . In addition, to avoid the confounding factor of the onlyCancer = TRUE argument, we will set it to FALSE, so we measure directly the time of individual runs.

#### 2.3.1 Exponential model with 10,000 and 50,000 genes

#### 2.3.1.1 Exponential, 10,000 genes, example 1

We will start with 10000 genes and an exponential model, where we stop when the population grows over  $10^6$  individuals:

```
t_e_10000
##
             system elapsed
      user
##
     4.368
             0.196
                      4.566
summary(e_10000)[, c(1:3, 8, 9)]
     NumClones TotalPopSize LargestClone FinalTime NumIter
## 1
          5017
                     1180528
                                    415116
                                                  143
                                                         7547
## 2
          3726
                                                         5746
                     1052061
                                    603612
                                                  131
## 3
          4532
                     1100721
                                    259510
                                                  132
                                                         6674
          4150
                     1283115
                                    829728
                                                   99
                                                         6646
## 4
## 5
          4430
                     1139185
                                    545958
                                                  146
                                                         6748
print(object.size(e_10000), units = "MB")
## 863.9 Mb
```

Each simulation takes about 1 second but note that the number of clones for most simulations is already over 4000 and that the size of the returned object is close to 1 GB (a more detailed explanation of where this 1 GB comes from is deferred until section 2.3.1.6).

#### 2.3.1.2 Exponential, 10,000 genes, example 2

We can decrease the size of the returned object if we use the keepEvery = NA argument (this setting was explained in detail in section 2.1):

```
t_e_10000b
##
      user
            system elapsed
##
     5.484
             0.100
                      5.585
summary(e_10000b)[, c(1:3, 8, 9)]
     NumClones TotalPopSize LargestClone FinalTime NumIter
##
## 1
          2465
                     1305094
                                    727989
                                                   91
                                                         6447
```

```
## 2
           2362
                      1070225
                                     400329
                                                   204
                                                           8345
## 3
           2530
                      1121164
                                     436721
                                                   135
                                                           8697
## 4
           2593
                      1206293
                                                   125
                                                           8149
                                     664494
## 5
           2655
                      1186994
                                     327835
                                                   191
                                                           8572
print(object.size(e 10000b), units = "MB")
## 488.3 Mb
```

#### 2.3.1.3 Exponential, 50,000 genes, example 1

Let's use 50,000 genes. To keep object sizes reasonable we use keepEvery = NA. For now, we also set mutationPropGrowth = FALSE so that the mutation rate does not become really large in clones with many mutations but, of course, whether or not this is a reasonable decision depends on the problem; see also below.

```
ng < -50000
u <- allFitnessEffects(noIntGenes = c(rep(0.1, ng/2),
                                       rep(-0.1, ng/2)))
t_e_50000 <- system.time(
    e_50000 <- oncoSimulPop(5,</pre>
                             model = "Exp",
                             mu = 1e-7,
                             detectionSize = 1e6,
                             detectionDrivers = NA,
                             detectionProb = NA,
                             keepPhylog = TRUE,
                             onlyCancer = FALSE,
                             keepEvery = NA,
                             mutationPropGrowth = FALSE,
                             mc.cores = 1
                             ))
t e 50000
##
      user
            system elapsed
    44.192
            1.684 45.891
summary(e_50000)[, c(1:3, 8, 9)]
     NumClones TotalPopSize LargestClone FinalTime NumIter
##
## 1
          7367
                     1009949
                                                75.00
                                    335455
                                                        18214
## 2
                                                63.65
                                                        17379
          8123
                     1302324
                                    488469
## 3
                                                72.57
                                                        21144
          8408
                     1127261
                                    270690
                     1138513
                                    318152
                                                80.59
                                                        20994
## 4
          8274
## 5
          7520
                     1073131
                                    690814
                                                70.00
                                                        18569
```

```
print(object.size(e_50000), units = "MB")
## 7598.6 Mb
```

Of course, simulations now take longer and the size of the returned object is over 7 GB (we are keeping more than 7,000 clones, even if when we prune all those that went extinct).

#### 2.3.1.4 Exponential, 50,000 genes, example 2

What if we had not pruned?

```
ng <- 50000
u <- allFitnessEffects(noIntGenes = c(rep(0.1, ng/2),
                                     rep(-0.1, ng/2))
t_e_50000np <- system.time(
   e 50000np <- oncoSimulPop(5,
                             model = "Exp",
                             mu = 1e-7,
                             detectionSize = 1e6,
                             detectionDrivers = NA,
                             detectionProb = NA,
                             keepPhylog = TRUE,
                             onlyCancer = FALSE,
                             keepEvery = 1,
                             mutationPropGrowth = FALSE,
                             mc.cores = 1
                             ))
t e 50000np
## user system elapsed
          2.764 45.079
## 42.316
summary(e_50000np)[, c(1:3, 8, 9)]
##
     NumClones TotalPopSize LargestClone FinalTime NumIter
## 1
         13406
                   1027949
                                  410074
                                            71.97 19469
## 2
         12469
                    1071325
                                  291852
                                             66.00
                                                     17834
## 3
         11821
                    1089834
                                  245720
                                             90.00
                                                     16711
         14008
                                  505607
                                             77.61
## 4
                    1165168
                                                     19675
## 5
        14759
                    1074621
                                  205954
                                             87.68
                                                     20597
print(object.size(e_50000np), units = "MB")
## 12748.4 Mb
```

The main effect is not on execution time but on object size (it has grown by 5 GB). We are tracking more than 10,000 clones.

#### 2.3.1.5 Exponential, 50,000 genes, example 3

What about the mutationPropGrowth setting? We will rerun the example in 2.3.1.3 leaving keepEvery = NA but with the default mutationPropGrowth:

```
ng <- 50000
u <- allFitnessEffects(noIntGenes = c(rep(0.1, ng/2),
                                       rep(-0.1, ng/2))
t e 50000c <- system.time(
    e 50000c <- oncoSimulPop(5,
                             model = "Exp",
                             mu = 1e-7,
                             detectionSize = 1e6,
                             detectionDrivers = NA,
                             detectionProb = NA,
                             keepPhylog = TRUE,
                             onlyCancer = FALSE,
                             keepEvery = NA,
                             mutationPropGrowth = TRUE,
                             mc.cores = 1
                             ))
t_e_50000c
##
      user system elapsed
## 84.228
            2.416 86.665
summary(e 50000c)[, c(1:3, 8, 9)]
     NumClones TotalPopSize LargestClone FinalTime NumIter
## 1
         11178
                     1241970
                                   344479
                                               84.74
                                                       27137
## 2
                     1307086
         12820
                                   203544
                                               91.94
                                                       33448
## 3
         10592
                     1126091
                                   161057
                                               83.81
                                                       26064
                                   148986
## 4
         11883
                     1351114
                                               65.68
                                                       25396
         10518
                                   253523
                                               99.79
## 5
                     1101392
                                                       26082
print(object.size(e_50000c), units = "MB")
## 10904.9 Mb
```

As expected (because the mutation rate per unit time is increasing in the fastest growing clones), we have many more clones, larger objects, and longer times of execution here: we almost double the time and the size of the object increases by almost 3 GB.

What about larger population sizes or larger mutation rates? The number of clones starts growing fast, which means much slower execution times and much larger returned objects (see also the examples below).

#### 2.3.1.6 Interlude: where is that 1 GB coming from?

In section 2.3.1.1 we have seen an apparently innocuous simulation producing a returned object of almost 1 GB. Where is that coming from? It means that each simulation produced almost 200 MB of output.

Let us look at one simulation in more detail:

```
r1 <- oncoSimulIndiv(u,
                     model = "Exp",
                     mu = 1e-7,
                      detectionSize = 1e6,
                      detectionDrivers = NA,
                      detectionProb = NA,
                      keepPhylog = TRUE,
                      onlyCancer = FALSE,
                     mutationPropGrowth = TRUE
summary(r1)[c(1, 8)]
     NumClones FinalTime
## 1
          3887
                       345
print(object.size(r1), units = "MB")
## 160 Mb
## Size of the two largest objects inside:
sizes \leftarrow lapply(r1, function(x) object.size(x)/(1024^2))
sort(unlist(sizes), decreasing = TRUE)[1:2]
## Genotypes pops.by.time
         148.28
                        10.26
##
dim(r1$Genotypes)
## [1] 10000 3887
```

The above shows the reason: the **Genotypes** matrix is a 10,000 by 3,887 integer matrix (with a 0 and 1 indicating not-mutated/mutated for each gene in each genotype) and in R integers use 4 bytes each. The pops.by.time matrix is 346 by 3,888 (the 1 in 346 = 345 + 1 comes from starting at 0 and going up to the final time, both included; the 1 in 3888 = 3887 + 1 is from the column of time) double matrix and doubles use 8 bytes<sup>4</sup>.

#### 2.3.2 McFarland model with 50,000 genes; the effect of keepEvery

We show an example of McFarland's model with 50,000 genes in section 6.5.3. We will show here a few more examples with those many genes but with a different fitness specification and changing several other settings.

<sup>&</sup>lt;sup>4</sup>These matrices do not exist during most of the execution of the C++ code; they are generated right before returning from the C++ code.

#### 2.3.2.1 McFarland, 50,000 genes, example 1

Let's start with mutationPropGrowth = FALSE and keepEvery = NA. Simulations end when population size  $\geq 10^6$ .

```
ng <- 50000
u <- allFitnessEffects(noIntGenes = c(rep(0.1, ng/2),
                                    rep(-0.1, ng/2))
t mc 50000 nmpg <- system.time(
   mc 50000 nmpg <- oncoSimulPop(5,
                                model = "McFL",
                                mu = 1e-7,
                                detectionSize = 1e6,
                                detectionDrivers = NA,
                                detectionProb = NA,
                                keepPhylog = TRUE,
                                onlyCancer = FALSE,
                                keepEvery = NA,
                                mutationPropGrowth = FALSE,
                                mc.cores = 1
                                ))
t_mc_50000_nmpg
## user system elapsed
## 30.46
           0.54
                   31.01
summary(mc 50000 nmpg)[, c(1:3, 8, 9)]
##
    \it NumClones\ TotalPopSize\ LargestClone\ FinalTime\ NumIter
## 1
         1902
                  1002528
                              582752
                                           284.2 31137
## 2
        2159
                                           274.8 36905
                  1002679
                                404858
                                            334.5 42429
## 3
         2247
                  1002722
                                 185678
## 4
         2038
                  1009606
                                 493574
                                           218.4 32519
## 5
         2222
                   1004661
                                 162628
                                           291.0
                                                  38470
print(object.size(mc 50000 nmpg), units = "MB")
## 2057.6 Mb
```

We are already dealing with 2000 clones.

#### 2.3.2.2 McFarland, 50,000 genes, example 2

Setting keepEvery = 1 (i.e., keeping track of clones with an interval of 1):

```
model = "McFL",
                                    mu = 1e-7,
                                    detectionSize = 1e6,
                                    detectionDrivers = NA,
                                    detectionProb = NA,
                                    keepPhylog = TRUE,
                                    onlyCancer = FALSE,
                                    keepEvery = 1,
                                    mutationPropGrowth = FALSE,
                                    mc.cores = 1
                                    ))
t_mc_50000_nmpg_k
##
      user system elapsed
##
    30.000
            1.712 31.714
summary(mc_50000_nmpg_k)[, c(1:3, 8, 9)]
##
     NumClones TotalPopSize LargestClone FinalTime NumIter
## 1
          8779
                    1000223
                                   136453
                                              306.7
                                                      38102
## 2
                                              345.3
          7442
                    1006563
                                   428150
                                                      35139
## 3
          8710
                    1003509
                                   224543
                                              252.3
                                                      35659
## 4
          8554
                    1002537
                                   103889
                                              273.7
                                                      36783
## 5
          8233
                    1003171
                                   263005
                                              301.8
                                                      35236
print(object.size(mc 50000 nmpg k), units = "MB")
## 8101.4 Mb
```

Computing time increases slightly but the major effect is seen on the size of the returned object, that increases by a factor of about 4x, up to 8 GB, corresponding to the increase in about 4x in the number of clones being tracked (see details of where the size of this object comes from in section 2.3.1.6).

#### 2.3.2.3 McFarland, 50,000 genes, example 3

We will set keepEvery = NA again, but we will now increase detection size by a factor of 3 (so we stop when total population size becomes  $> 3 * 10^6$ ).

```
detectionDrivers = NA,
                                    detectionProb = NA,
                                    keepPhylog = TRUE,
                                    onlyCancer = FALSE,
                                    keepEvery = NA,
                                    mutationPropGrowth = FALSE,
                                    mc.cores = 1
                                    ))
t_mc_50000_nmpg_3e6
   user system elapsed
## 77.240 1.064 78.308
summary(mc 50000 nmpg 3e6)[, c(1:3, 8, 9)]
    \it NumClones\ TotalPopSize\ LargestClone\ FinalTime\ NumIter
## 1
         5487
                  3019083
                                836793 304.5 65121
                   3011816
## 2
         4812
                                 789146
                                           286.3 53087
## 3
                                1970957
                                           236.6 45918
         4463
                   3016896
## 4
         5045
                   3028142
                                956026
                                          360.3 63464
## 5
         4791
                                           358.1 55012
                   3029720
                                 916692
print(object.size(mc_50000_nmpg_3e6), units = "MB")
## 4759.3 Mb
```

Compared with the first run (2.3.2.1) we have approximately doubled computing time, number of iterations, number of clones, and object size.

#### 2.3.2.4 McFarland, 50,000 genes, example 4

Let us use the same detectionSize = 1e6 as in the first example (2.3.2.1), but with 5x the mutation rate:

```
t_mc_50000_nmpg_5mu
      user system elapsed
## 167.332
            1.796 169.167
summary(mc_50000_nmpg_5mu)[, c(1:3, 8, 9)]
     NumClones TotalPopSize LargestClone FinalTime NumIter
## 1
          7963
                    1004415
                                   408352
                                              99.03
                                                       57548
## 2
          8905
                    1010751
                                             130.30
                                                       74738
                                   120155
## 3
          8194
                    1005465
                                   274661
                                              96.98
                                                       58546
## 4
          9053
                    1014049
                                   119943
                                                       75379
                                              112.23
## 5
          8982
                    1011817
                                    95047
                                              99.95
                                                       76757
print(object.size(mc 50000 nmpg 5mu), units = "MB")
## 8314.4 Mb
```

The number of clones we are tracking is about 4x the number of clones of the first example (2.3.2.1), and roughly similar to the number of clones of the second example (2.3.2.2), and size of the returned object is similar to that of the second example. But computing time has increased by a factor of about 5x and iterations have increased by a factor of about 2x. Iterations increase because mutation is more frequent; in addition, at each sampling period each iteration needs to do more work as it needs to loop over a larger number of clones and this larger number includes clones that are not shown here, because they are pruned (they are extinct by the time we exit the simulation —again, pruning is discussed with further details in 14.2.1).

#### 2.3.2.5 McFarland, 50,000 genes, example 5

Now let's run the above example but with keepEvery = 1:

```
t mc 50000 nmpg 5mu k <- system.time(
    mc_50000_nmpg_5mu_k <- oncoSimulPop(5,
                                         model = "McFL",
                                        mu = 5e-7,
                                         detectionSize = 1e6,
                                         detectionDrivers = NA,
                                         detectionProb = NA,
                                         keepPhylog = TRUE,
                                         onlyCancer = FALSE,
                                         keepEvery = 1,
                                         mutationPropGrowth = FALSE,
                                         mc.cores = 1
                                         ))
t_mc_50000_nmpg 5mu k
      user system elapsed
## 174.404
           5.068 179.481
```

```
summary(mc 50000 nmpg 5mu k)[, c(1:3, 8, 9)]
##
     NumClones TotalPopSize LargestClone FinalTime NumIter
                                               123.4
## 1
         25294
                     1001597
                                    102766
                                                        74524
## 2
         23766
                                               124.3
                                                       71808
                     1006679
                                    223010
## 3
         21755
                                               114.8
                     1001379
                                    203638
                                                        62609
         24889
                     1012103
                                    161003
                                               119.3
                                                       75031
## 4
## 5
         21844
                     1002927
                                    255388
                                               108.8
                                                       64556
print(object.size(mc 50000 nmpg 5mu k), units = "MB")
## 22645.8 Mb
```

We have already seen these effects before in section 2.3.2.2: using keepEvery = 1 leads to a slight increase in execution time. What is really affected is the size of the returned object which increases by a factor of about 3x (and is now over 20GB). That 3x corresponds, of course, to the increase in the number of clones being tracked (now over 20,000). This, by the way, also allows us to understand the comment above, where we said that in these two cases (where we have increased mutation rate) at each iteration we need to do more work as at every update of the population the algorithm needs to loop over a much larger number of clones (even if many of those are eventually pruned).

#### 2.3.2.6 McFarland, 50,000 genes, example 6

Finally, we will run the example in section 2.3.2.1 with the default of mutationPropGrowth = TRUE:

```
t_mc_50000 <- system.time(
    mc 50000 <- oncoSimulPop(5,
                             model = "McFL",
                             mu = 1e-7,
                              detectionSize = 1e6,
                              detectionDrivers = NA,
                              detectionProb = NA,
                              keepPhylog = TRUE,
                              onlyCancer = FALSE,
                              keepEvery = NA,
                              mutationPropGrowth = TRUE,
                              mc.cores = 1
                              ))
t mc 50000
      user
            system elapsed
            2.808 306.223
## 303.352
```

```
summary(mc_50000)[, c(1:3, 8, 9)]
     NumClones TotalPopSize LargestClone FinalTime NumIter
##
## 1
         13928
                     1010815
                                   219814
                                               210.9
                                                        91255
## 2
                                               178.1
         12243
                     1003267
                                   214189
                                                        67673
## 3
         13880
                     1014131
                                    124354
                                               161.4
                                                        88322
                                     75521
                                               205.7
## 4
         14104
                     1012941
                                                        98583
## 5
         12428
                     1005594
                                    232603
                                               167.4
                                                        70359
print(object.size(mc 50000), units = "MB")
## 12816.6 Mb
```

Note the huge increase in computing time (related of course to the huge increase in number of iterations) and in the size of the returned object: we have gone from having to track about 2000 clones to tracking over 12000 clones even when we prune all clones without descendants.

#### **2.3.3** Examples with s = 0.05

A script with the above runs but using s = 0.05 instead of s = 0.1 is available from the repository ('miscell-files/vignette\_bench\_Rout/large\_num\_genes\_0.05.Rout'). I will single out a couple of cases here.

First, we repeat the run shown in section 2.3.2.5:

```
t_mc_50000_nmpg_5mu_k <- system.time(
    mc 50000 nmpg 5mu k <- oncoSimulPop(2,
                                        u,
                                        model = "McFL",
                                        mu = 5e-7,
                                        detectionSize = 1e6,
                                        detectionDrivers = NA,
                                        detectionProb = NA,
                                        keepPhylog = TRUE,
                                        onlyCancer = FALSE,
                                        keepEvery = 1,
                                        mutationPropGrowth = FALSE,
                                        mc.cores = 1
                                        ))
t mc 50000 nmpg 5mu k
      user system elapsed
## 305.512
             5.164 310.711
summary(mc_50000_nmpg_5mu_k)[, c(1:3, 8, 9)]
##
     NumClones TotalPopSize LargestClone FinalTime NumIter
## 1
                    1003273
         61737
                                   104460 295.8731
                                                      204214
## 2
         65072
                    1000540
                                   133068 296.6243 210231
```

```
print(object.size(mc_50000_nmpg_5mu_k), units = "MB")
## 24663.6 Mb
```

Note we use only two replicates, since those two already lead to a 24 GB returned object as we are tracking more than 60,000 clones, more than twice those with s=0.1. The reason for the difference in number of clones and iterations is of course the change from s=0.1 to s=0.05: under the McFarland model to reach population sizes of  $10^6$  starting from an equilibrium population of 500 we need about 43 mutations (whereas only about 22 are needed if  $s=0.1^5$ ).

Next, let us rerun 2.3.2.1:

```
t_mc_50000_nmpg <- system.time(
    mc 50000 nmpg <- oncoSimulPop(5,
                                   model = "McFL",
                                   mu = 1e-7,
                                   detectionSize = 1e6,
                                   detectionDrivers = NA,
                                   detectionProb = NA,
                                   keepPhylog = TRUE,
                                   onlyCancer = FALSE,
                                   keepEvery = NA,
                                   mutationPropGrowth = FALSE,
                                   mc.cores = 1
                                   ))
t mc 50000 nmpg
      user system elapsed
## 111.236
             0.596 111.834
summary(mc_50000_nmpg)[, c(1:3, 8, 9)]
     NumClones TotalPopSize LargestClone FinalTime NumIter
## 1
                     1000700
          2646
                                   217188
                                             734.475
                                                      108566
## 2
          2581
                     1001626
                                   209873
                                             806.500
                                                      107296
## 3
          2903
                     1001409
                                   125148
                                             841.700
                                                      120859
                                   473948
## 4
          2310
                     1000146
                                             906.300
                                                       91519
## 5
          2704
                     1001290
                                   448409
                                             838.800
                                                      103556
print(object.size(mc 50000 nmpg), units = "MB")
## 2638.3 Mb
```

Using s=0.05 leads to a large increase in final time and number of iterations. However, as we are using the keepEvery = NA setting, the increase in number of

<sup>&</sup>lt;sup>5</sup>Given the dependence of death rates on population size in McFarland's model (section 3.2.1 and 3.2.1.1), if all mutations have the same fitness effects we can calculate the equilibrium population size (where birth and death rates are equal) for a given number of mutated genes as:  $K*(e^{(1+s)^p}-1)$ , where K is the initial equilibrium size, s the fitness effect of each mutation, and p the number of mutated genes.

clones tracked and in size of returned object is relatively small.

## 2.3.4 The different consequences of keepEvery = NA in the Exp and McFL models

We have seen that keepEvery = NA often leads to much smaller returned objects when using the McFarland model than when using the Exp model. Why? Because in the McFarland model there is strong competition and there can be complete clonal sweeps so that in extreme cases a single clone might be all that is left after some time. This is not the case in the exponential models.

Of course, the details depend on the difference in fitness effects between different genotypes (or clones). In particular, we have seen several examples where even with keepEvery=NA there are a lot of clones in the McFL models. In those examples many clones had identical fitness (the fitness effects of all genes with positive fitness was the same, and ditto for the genes with negative fitness effects), so no clone ends up displacing all the others.

#### 2.3.5 Are we keeping the complete history (genealogy) of the clones?

Yes we are if we run with keepPhylog = TRUE, regardless of the setting for keepEvery. As explained in section 14.2, OncoSimulR prunes clones that never had a population size larger than zero at any sampling period (so they are not reflected in the pops.by.time matrix in the output). And when we set keepEvery = NA we are telling OncoSimulR to discard all sampling periods except the very last one (i.e., the pops.by.time matrix contains only the clones with 1 or more cells at the end of the simulation).

keepPhylog operates differently: it records the exact time at which a clone appeared and the clone that gave rise to it. This information is kept regardless of whether or not those clones appear in the pops.by.time matrix.

Keeping the complete genealogy might be of limited use if the pops.by.time matrix only contains the very last period. However, you can use plotClonePhylog and ask to be shown only clones that exist in the very last period (while of course showing all of their ancestors, even if those are now extinct —i.e., regardless of their abundance).

For instance, in run 2.3.1.3 we could have looked at the information stored about the genealogy of clones by doing (we look at the first "individual" of the simulation, of the five "individuals" we simulated):

```
head(e_50000[[1]]$other$PhylogDF)
     parent child
##
                     time
## 1
             3679 0.8402
             4754 1.1815
## 2
## 3
            20617 1.4543
## 4
            15482 2.3064
             4431 3.7130
## 5
## 6
            41915 4.0628
```

```
tail(e 50000[[1]]$other$PhylogDF)
##
                             parent
                                                                child time
## 20672
                        3679, 20282
                                                   3679, 20282, 22359 75.0
## 20673
                3679, 17922, 22346
                                            3679, 17922, 22346, 35811 75.0
                                                    2142, 3679, 25838 75.0
## 20674
                         2142, 3679
## 20675
                3679, 17922, 19561
                                            3679, 17922, 19561, 43777 75.0
## 20676 3679, 15928, 19190, 20282 3679, 15928, 19190, 20282, 49686 75.0
## 20677
                 2142, 3679, 16275
                                             2142, 3679, 16275, 24201 75.0
```

where each row corresponds to one event of appearance of a new clone, the column labeled "parent" are the mutated genes in the parent, and the column labeled "child" are the mutated genes in the child.

And we could plot the genealogical relationships of clones that have a population size of at least one in the last period (again, while of course showing all of their ancestors, even if those are now extinct —i.e., regardless of their current numbers) doing:

```
plotClonePhylog(e_50000[[1]]) ## plot not shown
```

What is the cost of keep the clone genealogies? In terms of time it is minor. In terms of space, and as shown in the example above, we can end up storing a data frame with tends of thousands of rows and three columns (two factors, one float). In the example above the size of that data frame is approximately 2 MB for a single simulation. This is much smaller than the pops.by.time or Genotypes matrices, but it can quickly build up if you routinely launch, say, 1000 simulations via oncoSimulPop. That is why the default is keepPhylog = FALSE as this information is not needed as often as that in the other two matrices (pops.by.time and Genotypes).

## 2.4 Population sizes $\geq 10^{10}$

We have already seen examples where population sizes reach  $10^8$  to  $10^{10}$ , as in Tables 3, 5, 7. What about even larger population sizes?

The C++ code will unconditionally alert if population sizes exceed  $4 * 10^{15}$  as in those cases loosing precision (as we are using doubles) would be unavoidable, and we would also run into problems with the generation of binomial random variates (code that illustrates and discusses this problem is available in file "example-binom-problems.cpp", in directory "/inst/miscell"). However, well before we reach  $4 * 10^{15}$  we loose precision from other sources. One of the most noticeable ones is that when we reach population sizes around  $10^{11}$  the C++ code will often alert us by throwing exceptions with the message Recoverable exception ti set to DBL\_MIN. Rerunning. I throw this exception because  $t_i$ , the random variable for time to next mutation, is less than DBL\_MIN, the minimum representable floating-point number. This happens because, unless we use really tiny mutation rates, the time to a mutation starts getting closer to zero as population sizes grow very large. It might be possible to ameliorate these problems somewhat by using long doubles (instead of doubles) or special purpose libraries that provide more precision. However,

this would make it harder to run the same code in different operating systems and would likely decrease execution speed on the rest of the common scenarios for which OncoSimulR has been designed.

The following code shows some examples where we use population sizes of  $10^{10}$  or larger. Since we do not want simulations in the exponential model to end because of extinction, I use a fitness specification where all genes have a positive fitness effect and we start all simulations from a large population (to make it unlikely that the population will become extinct before cells mutate and start increasing in numbers). We set the maximum running time to 10 minutes. We keep the genealogy of the clones and use keepEvery = 1.

```
ng <- 50
u <- allFitnessEffects(noIntGenes = c(rep(0.1, ng)))</pre>
```

```
t_mc_k_50_1e11 <- system.time(
   mc k 50 1e11 <- oncoSimulPop(5,
                                u,
                                model = "McFL",
                                mu = 1e-7,
                                detectionSize = 1e11,
                                initSize = 1e5,
                                detectionDrivers = NA,
                                detectionProb = NA,
                                keepPhylog = TRUE,
                                onlyCancer = FALSE,
                                mutationPropGrowth = FALSE,
                                keepEvery = 1,
                                finalTime = 5000,
                                mc.cores = 1,
                                max.wall.time = 600
                                ))
## Recoverable exception ti set to DBL MIN. Rerunning.
## Recoverable exception ti set to DBL MIN. Rerunning.
t_mc_k_50_1e11
## user system elapsed
## 613.612 0.040 613.664
summary(mc_k_50_1e11)[, c(1:3, 8, 9)]
    NumClones TotalPopSize LargestClone FinalTime NumIter
## 1
          5491 100328847809 44397848771 1019.950 942764
## 2
          3194 100048090441 34834178374 789.675
                                                    888819
## 3
          5745 100054219162 24412502660
                                           927.950
                                                    929231
          4017 101641197799 60932177160 750.725 480938
## 4
## 5
         5393 100168156804 41659212367 846.250 898245
```

```
## print(object.size(mc_k_50_1e11), units = "MB")
## 177.8 Mb
```

We get to  $10^{11}$ . But notice the exception with the warning about  $t_i$ . Notice also that this takes a long time and we run a very large number of iterations (getting close to one million in some cases).

Now the exponential model with detectionSize = 1e11:

```
t exp k 50 1e11 <- system.time(
    exp k 50 1e11 <- oncoSimulPop(5,
                                  model = "Exp",
                                  mu = 1e-7.
                                  detectionSize = 1e11,
                                  initSize = 1e5,
                                  detectionDrivers = NA,
                                  detectionProb = NA,
                                  keepPhylog = TRUE,
                                  onlyCancer = FALSE,
                                  mutationPropGrowth = FALSE,
                                  keepEvery = 1,
                                  finalTime = 5000,
                                  mc.cores = 1,
                                  max.wall.time = 600,
                                  errorHitWallTime = FALSE,
                                  errorHitMaxTries = FALSE
                                  ))
## Recoverable exception ti set to DBL_MIN. Rerunning.
## Hitted wall time. Exiting.
## Recoverable exception ti set to DBL MIN. Rerunning.
## Recoverable exception ti set to DBL MIN. Rerunning.
## Recoverable exception ti set to DBL_MIN. Rerunning.
## Hitted wall time. Exiting.
## Recoverable exception ti set to DBL MIN. Rerunning.
## Recoverable exception ti set to DBL_MIN. Rerunning.
## Recoverable exception ti set to DBL_MIN. Rerunning.
## Recoverable exception ti set to DBL MIN. Rerunning.
## Recoverable exception ti set to DBL_MIN. Rerunning.
## Hitted wall time. Exiting.
## Hitted wall time. Exiting.
t_exp_k_50_1e11
              system elapsed
##
       user
## 2959.068
              0.128 2959.556
try(summary(exp_k_50_1e11)[, c(1:3, 8, 9)])
```

```
##
    NumClones TotalPopSize LargestClone FinalTime NumIter
## 1
          6078 65172752616
                            16529682757
                                          235.7590 1883438
## 2
          5370 106476643712 24662446729
                                          232.0000 2516675
## 3
          2711 21911284363
                            17945303353
                                          224.8608
                                                    543698
          2838 13241462284
                              2944300245
                                          216.8091
                                                    372298
## 4
                76166784312
                            10941729810
## 5
          7289
                                          240.0217 1999489
print(object.size(exp k 50 1e11), units = "MB")
## 53.5 Mb
```

Note that we almost reached max.wall.time (600 \* 5 = 3000). What if we wanted to go up to  $10^{12}$ ? We would not be able to do it in 10 minutes. We could set max.wall.time to a value larger than 600 to allow us to reach larger sizes but then we would be waiting for a possibly unacceptable time for simulations to finish. Moreover, this would eventually fail as simulations would keep hitting the  $t_i$  exception without ever being able to complete. Finally, even if we were very patient, hitting that  $t_i$  exception should make us worry about possible biases in the samples.

# 2.5 A summary of some determinants of running time and space consumption

To summarize this section, we have seen:

- Both McFL and Exp can be run in short times over a range of sizes for the detectionProb and detectionSize mechanisms using a complex fitness specification with moderate numbers of genes. These are the typical or common use cases of OncoSimulR.
- The keepEvery argument can have a large effect on time in the McFL models and specially on object sizes. If only the end result of the simulation is to be used, you should set keepEvery = NA.
- The distribution of fitness effects and the fitness landscape can have large effects on running times. Sometimes these are intuitive and simple to reason about, sometimes they are not as they interact with other factors (e.g., stopping mechanism, numbers of clones, etc). In general, there can be complex interactions between different settings, from mutation rate to fitness effects to initial size. As usual, test before launching a massive simulation.
- Simulations start to slow down and lead to a very large object size when we keep track of around 6000 to 10000 clones. Anything that leads to these patterns will slow down the simulations.
- OncoSimulR needs to keep track of genotypes (or clones), not just numbers of drivers and passengers, because it allows you to use complex fitness and mutation specifications that depend on specific genotypes. The keepEvery = NA is an approach to store only the minimal information needed, but it is unavoidable that during the simulations we might be forced to deal with many thousands of different clones.

## 3 Specifying fitness effects

OncoSimulR uses a standard continuous time model, where individual cells divide, die, and mutate with rates that can depend on genotype and population size; over time the abundance of the different genotypes changes by the action of selection (due to differences in net growth rates among genotypes), drift, and mutation. As a result of a mutation in a pre-existing clone new clones arise, and the birth rate of a newly arisen clone is determined at the time of its emergence as a function of its genotype. Simulations can use an use exponential growth model or a model with carrying capacity that follows McFarland et al. (2013). For the exponential growth model, the death rate is fixed at one whereas in the model with carrying capacity death rate increases with population size. In both cases, therefore, fitness differences among genotypes in a given population at a given time are due to differences in the mapping between genotype and birth rate. There is second exponential model (called "Bozic") where birth rate is fixed at one, and genotype determines death rate instead of birth rate (see details in 3.2). So when we discuss specifying fitness effects or the effects of genes on fitness, we are actually referring to specifying effects on birth (or death) rates, which then translate into differences in fitness (since the other rate, death or birth, is either fixed, as in the Exp and Bozic models, or depends on the population size). This is also shown in Table 1, in the rows for "Fitness components", under "Evolutionary Features".

In the case of frequency-dependent fitness simulations (10), the fitness effects must be reevaluated frequently so that birth rate, death rate, or both, depending the model used, are updated. To do this it is necessary to use a short step to reevaluate fitness; this is done using a small value for sampleEvery parameter in oncoSimulindv (see 14.8 for more details), as is the case when using McFarland model.

Incidentally, notice that with OncoSimulR we do not directly specify fitness itself (even if, for the sake of simplicity, we often refer to fitness in the documentation) as fitness is, arguably, a derived quantity (Doebeli, Ispolatov, and Simon 2017). Rather, we specify how birth and/or death rates, which are the actual mechanistic drivers of evolutionary dynamics, are related to genotypes (or to the frequencies of the different genotypes).

### 3.1 Introduction to the specification of fitness effects

With OncoSimulR you can specify different types of effects on fitness:

• A special type of epistatic effect that is particularly amenable to be represented as a graph (a DAG). In this graph having, say, "B" be a child of "A" means that a mutation in B can only accumulate if a mutation in A is already present. This is what OT (Desper et al. 1999; Szabo and Boucher 2008), CBN (Beerenwinkel, Eriksson, and Sturmfels 2007; Gerstung et al. 2009, 2011), progression networks (Farahani and Lagergren 2013), and other similar models (Korsunsky et al. 2014) generally mean. Details are provided in section 3.4. Note that this is not an order effect (discussed below): the fitness of a genotype from this DAGs is a function of whether or not the restrictions in the graph are satisfied, not the

historical sequence of how they were satisfied.

- Effects where the order in which mutations are acquired matters, as illustrated in section 3.6. There is, in fact, empirical evidence of these effects (Ortmann et al. 2015). For instance, the fitness of genotype "A, B" would differ depending on whether A or B was acquired first (or, as in the actual example in (Ortmann et al. 2015), the fitness of the mutant with JAK2 and TET2 mutated will depend on which of the genes was mutated first).
- General epistatic effects (e.g., section 3.7), including synthetic viability (e.g., section 3.9) and synthetic lethality/mortality (e.g., section 3.10).
- Genes that have independent effects on fitness (section 3.3).
- Modules (see section 3.5) allow you to specify any of the above effects (except those for genes without interactions, as it would not make sense there) in terms of modules (sets of genes), not individual genes. We will introduce them right after 3.4, and we will continue using them thereafter.

A guiding design principle of OncoSimulR is to try to make the specification of those effects as simple as possible but also as flexible as possible. Thus, there are two main ways of specifying fitness effects:

- Combining different types of effects in a single specification. For instance, you can combine epistasis with order effects with no interaction genes with modules. What you would do here is specify the effects that different mutations (or their combinations) have on fitness (the fitness effects) and then have OncoSimulR take care of combining them as if each of these were lego pieces. We will refer to this as the **lego system of fitness effects**. (As explained above, I find this an intuitive and very graphical analogy, which I have copied from Hothorn et al. (2006) and Hothorn et al. (2008)).
- Explicitly passing to OncoSimulR a mapping of genotypes to fitness. Here you specify the fitness of each genotype. We will refer to this as the **explicit mapping of genotypes to fitness**.

Both approaches have advantages and disadvantages. Here I emphasize some relevant differences.

- With the lego system you can specify huge genomes with an enormous variety of interactions, since the possible genotypes are not constructed in advance. You would not be able to do this with the explicit mapping of genotypes to fitness if you wanted to, say, construct that mapping for a modest genotype of 500 genes (you'd have more genotypes than particles in the observable Universe).
- For many models/data you often intuitively start with the fitness of the genotypes, not the fitness consequences of the different mutations. In these cases, you'd need to do the math to specify the terms you want if you used the lego system so you'll probably use the specification with the direct mapping genotype → fitness.

- Likewise, sometimes you already have a moderate size genotype → fitness mapping and you certainly do not want to do the math by hand: here the lego system would be painful to use.
- But sometimes we do think in terms of "the effects on fitness of such and such mutations are" and that immediately calls for the lego system, where you focus on the effects, and let OncoSimulR take care of doing the math of combining.
- If you want to use order effects, you must use the lego system (at least for now).
- If you want to specify modules, you must use the lego system (the explicit mapping of genotypes is, by its very nature, ill-suited for this).
- The lego system might help you see what your model really means: in many cases, you can obtain fairly succinct specifications of complex fitness models with just a few terms. Similarly, depending on what your emphasis is, you can often specify the same fitness landscape in several different ways.

Regardless of the route, you need to get that information into OncoSimulR's functions. The main function we will use is allFitnessEffects: this is the function in charge of reading the fitness specifications. We also need to discuss how, what, and where you have to pass to allFitnessEffects.

## 3.1.1 Explicit mapping of genotypes to fitness

Conceptually, the simplest way to specify fitness is to specify the mapping of all genotypes to fitness explicitly. An example will make this clear. Let's suppose you have a simple two-gene scenario, so a total of four genotypes, and you have a data frame with genotypes and fitness, where genoytpes are specified as character vectors, with mutated genes separated by commas:

```
m4 \leftarrow data.frame(G = c("WT", "A", "B", "A, B"), F = c(1, 2, 3, 4))
```

Now, let's give that to the allFitnessEffects function:

```
fem4 <- allFitnessEffects(genotFitness = m4)
## Warning in to_genotFitness_std(genotFitness,
## frequencyDependentFitness = FALSE, : First column of genotype
## fitness is a factor. Converting to character.
## Column names of object not Genotype and Fitness. Renaming them assuming that i</pre>
```

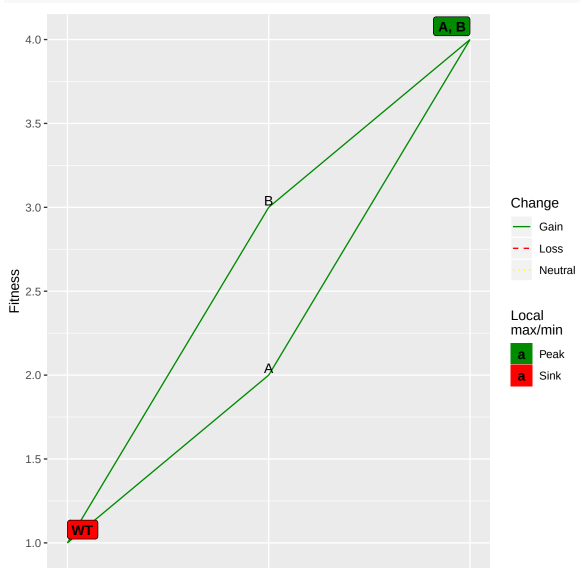
(The message is just telling you what the program guessed you wanted.)

That's it. You can try to plot that fitnessEffects object

```
try(plot(fem4))
## Error in plot.fitnessEffects(fem4) :
## This fitnessEffects object can not be ploted this way. It is probably one wi
```

In this case, you probably want to plot the fitness landscape.

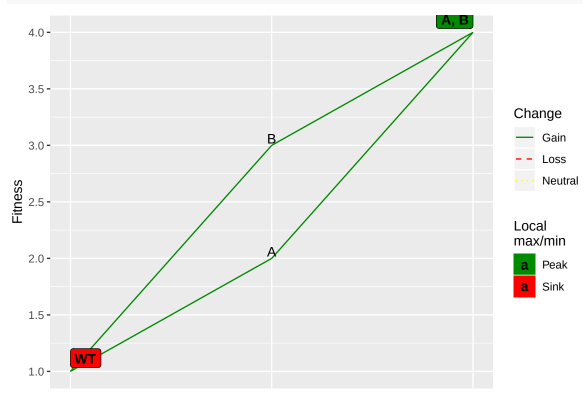




You can also check what OncoSimulR thinks the fitnesses are, with the evalAllGenotypes function that we will use repeatedly below (of course, here we should see the same fitnesses we entered):

And you can plot the fitness landscape:

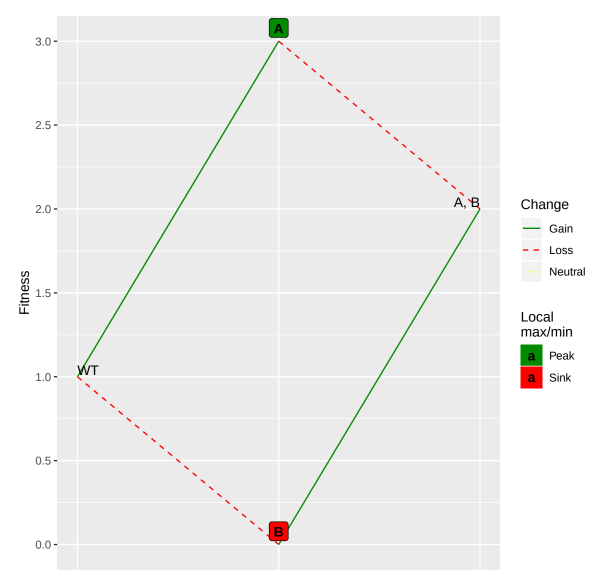




To specify the mapping you can also use a matrix (or data frame) with g+1 columns; each of the first g columns contains a 1 or a 0 indicating that the gene of that column is mutated or not. Column g+1 contains the fitness values. And you do not even need to specify all the genotypes: the missing genotypes are assigned a fitness 0—except for the WT genotype which, if missing, is assigned a fitness of 1:

```
m6 \leftarrow cbind(c(1, 1), c(1, 0), c(2, 3))
fem6 <- allFitnessEffects(genotFitness = m6)</pre>
## No column names: assigning gene names from LETTERS
## Warning in to_genotFitness_std(genotFitness,
\#\# frequency Dependent Fitness = FALSE, : No wild type in the fitness
## landscape!!! Adding it with fitness 1.
evalAllGenotypes(fem6, addwt = TRUE)
     Genotype Fitness
##
## 1
            WT
## 2
             A
                     3
## 3
             В
                     0
                     2
         A, B
## 4
## plot(fem6)
```

plotFitnessLandscape(evalAllGenotypes(fem6))



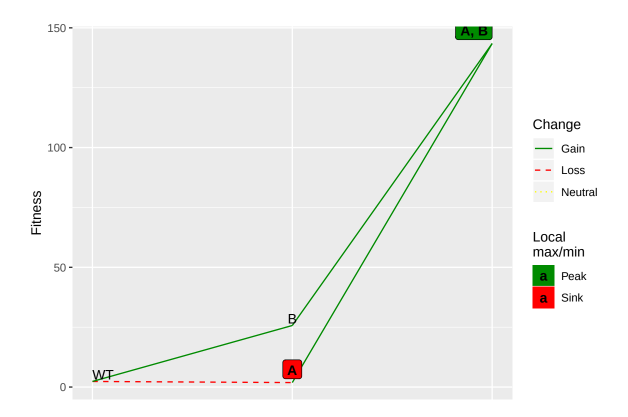
This way of giving a fitness specification to OncoSimulR might be ideal if you directly generate random mappings of genotypes to fitness (or random fitness landscapes), as we will do in section 9. Specially when the fitness landscape contains many non-viable genotypes (which are considered those with fitness —birth rate— < 1e - 9) this can result in considerable savings as we only need to check the fitness of the viable genotypes in a table (a C++ map). Note, however, that using the Bozic model with the fitness landscape specification is not tested. In addition, for speed, missing genotypes from the fitness landscape specification are taken to be non-viable genotypes (beware!! this is a breaking change relative to versions < 2.9.1)<sup>6</sup>.

In the case of frequency-dependent fitness situations, the only way to specify fitness effects is using genoFitnes as we have shown before, but now you need to set frequencyDependentFitness = TRUE in allFitnessEffects. The fundamental

<sup>&</sup>lt;sup>6</sup>Note for curious readers: it used to be the case that we converted the table of fitness of genotypes to a fitness specification with all possible epistatic interactions; you can take a look at the test file test.genot\_fitness\_to\_epistasis.R that uses the fem6 object. We no longer do that but instead pass directly the fitness landscape.

difference is the Fitness column in <code>genoFitnes</code>. Now this column must be a character vector and each element (character also) is a function whose variables are the relative frequencies of the clones in the population. You must specify the variables like f\_, for frequency of wild type, f\_1 or f\_A for frequency of mutant A or position 1, f\_1\_2 or f\_A\_B for double mutant, and so on. Mathematical operations and symbols allowed are described in the documentation of C++ library ExprTk (http://www.partow.net/programming/exprtk/). ExprTk is the library used to parse and evaluate the fitness equations. The numeric vector <code>spPopSizes</code> is only necesary to evaluate genotypes through <code>evalGenotype</code> or <code>evalAllGenotypes</code> functions because population sizes are needed to calculate the clone's frequencies.

```
r <- data.frame(Genotype = c("WT", "A", "B", "A, B"),
                 Fitness = c("10*f",
                             "10*f 1",
                             "50*f 2",
                             "200*(f 1 + f 2) + 50*f 1 2"))
afe <- allFitnessEffects(genotFitness = r,</pre>
                         frequencyDependentFitness = TRUE,
                         frequencyType = "rel",
                         spPopSizes = c(2500, 2000, 5500, 700))
## Warning in to_genotFitness_std(genotFitness,
## frequencyDependentFitness = TRUE, : First column of genotype fitness
## is a factor. Converting to character.
## Warning in to_genotFitness_std(genotFitness,
## frequencyDependentFitness = TRUE, : Second column of genotype
## fitness is a factor. Converting to character.
plotFitnessLandscape(evalAllGenotypes(afe))
```



#### 3.1.2 How to specify fitness effects with the lego system

An alternative general approach followed in many genetic simulators is to specify how particular combinations of alleles modify the wildtype genotype or the genotype that contains the individual effects of the interacting genes (e.g., see equation 1 in the supplementary material for FFPopSim (Zanini and Neher 2012)). For example, if we specify that a mutation in "A" contributes 0.04, a mutation in "B" contributes 0.03, and the double mutation "A:B" contributes 0.1, that means that the fitness of the "A, B" genotype (the genotype with A and B mutated) is that of the wildtype (1, by default), plus (actually, times —see section 3.2— but plus on the log scale) the effects of having A mutated, plus (times) the effects of having B mutated, plus (times) the effects of "A:B" both being mutated.

We will see below that with the "lego system" it is possible to do something very similar to the explicit mapping of section 3.1.1. But this will sometimes require a more cumbersome notation (and sometimes also will require your doing some math). We will see examples in sections 3.7.1, 3.7.2 and 3.7.3 or the example in 5.4.2. But then, if we can be explicit about (at least some of) the mappings  $genotype \rightarrow fitness$ , how are these procedures different? When you use the "lego system" you can combine both a partial explicit mapping of genotypes to fitness with arbitrary fitness effects of other genes/modules. In other words, with the "lego system" OncoSimulR makes it simple to be explicit about the mapping of specific genotypes, while also using the "how this specific effects modifies previous effects" logic, leading to a flexible specification. This also means that in many cases the same fitness effects can be specified in several different ways.

Most of the rest of this section is devoted to explaining how to combine those pieces.

Before that, however, we need to discuss the fitness model we use.

## 3.2 Numeric values of fitness effects

We evaluate fitness using the usual (Zanini and Neher 2012; Gillespie 1993; Beerenwinkel, Eriksson, and Sturmfels 2007; Datta et al. 2013) multiplicative model: fitness is  $\Pi(1+s_i)$  where  $s_i$  is the fitness effect of gene (or gene interaction) i. In all models except Bozic, this fitness refers to the growth rate (the death rate being fixed to  $1^7$ ). The original model of McFarland et al. (2013) has a slightly different parameterization, but you can go easily from one to the other (see section 3.2.1).

For the Bozic model (Bozic et al. 2010), however, the birth rate is set to 1, and the death rate then becomes  $\prod (1 - s_i)$ .

## 3.2.1 McFarland parameterization

In the original model of McFarland et al. (2013), the effects of drivers contribute to the numerator of the birth rate, and those of the (deleterious) passengers to the denominator as:  $\frac{(1+s)^d}{(1+s_p)^p}$ , where d and p are, respectively, the total number of drivers and passengers in a genotype, and here the fitness effects of all drivers is the same (s) and that of all passengers the same too  $(s_p)$ . Note that, as written above, and as explicitly said in McFarland et al. (2013) (see p. 2911) and CD McFarland (2014) (see p. 9), "(...)  $s_p$  is the fitness disadvantage conferred by a passenger". In other words, the larger the  $s_p$  the more deleterious the passenger.

This is obvious, but I make it explicit because in our parameterization a positive s means fitness advantage, whereas fitness disadvantages are associated with negative s. Of course, if you rewrite the above expression as  $\frac{(1+s)^d}{(1-s_p)^p}$  then we are back to the "positive means fitness advantage and negative means fitness disadvantage".

As CD McFarland (2014) explains (see p. 9, bottom), we can rewrite the above expression so that there are no terms in the denominator. McFarland writes it as (I copy verbatim from the fourth and fifth lines from the bottom on his p. 9)  $(1 + s_d)^{n_d} (1 - s_p')^{n_p}$  where  $s_p' = s_p/(1 + s_p)$ .

However, if we want to express everything as products (no ratios) and use the "positive s means advantage and negative s means disadvantage" rule, we want to write the above expression as  $(1 + s_d)^{n_d} (1 + s_{pp})^{n_p}$  where  $s_{pp} = -s_p/(1 + s_p)$ . And this is actually what we do in v.2. There is an example, for instance, in section 6.5.2 where you will see:

```
sp <- 1e-3
spp <- -sp/(1 + sp)
```

so we are going from the "(...)  $s_p$  is the fitness disadvantage conferred by a passenger" in McFarland et al. (2013) (p. 2911) and CD McFarland (2014) (p. 9) to the expression where we have a product  $\prod (1+s_i)$ , with the "positive s means advantage and negative s means disadvantage" rule. This reparameterization applies to v.2. In v.1 we use

<sup>&</sup>lt;sup>7</sup>You can change this if you really want to.

the same parameterization as in the original one in McFarland et al. (2013), but with the "positive s means advantage and negative s means disadvantage" rule (so we are using expression  $\frac{(1+s)^d}{(1-s_n)^p}$ ).

#### 3.2.1.1 Death rate under the McFarland model

For death rate, we use the expression that McFarland et al. (2013) (see their p. 2911) use "(...) for large cancers (grown to  $10^6$  cells)":  $D(N) = \log(1 + N/K)$  where K is the initial equilibrium population size. As the authors explain, for large N/K the above expression "(...) recapitulates Gompertzian dynamics observed experimentally for large tumors".

By default, OncoSimulR uses a value of  $K = initSize/(e^1 - 1)$  so that the starting population is at equilibrium.

A consequence of the above expression for death rate is that if the population size decreases the death rate decreases. This is not relevant in most cases (as mutations, or some mutations, will inexorably lead to population size increases). And this prevents the McFL model from resulting in extinction even with very small population sizes as long as birth rate  $\geq$  death rate. (For small population sizes, it is likely that the population will become extinct if birth rate = death rate; you can try this with the exponential model).

But this is not what we want in some other models, such as frequency-dependent ones, where modeling population collapse (which will happen if birth rate < death rate) can be important (as in the example in 10.3). Here, it makes sense to set  $D(N) = \max(1, \log(1 + N/K))$  so that the death rate never decreases below 1. (Using 1 is reasonable if we consider the equilibrium birth rate in the absence of any mutants to be 1). You can specify this behaviour using model McFLD (a shorthand for McFarlandLogD).

#### 3.2.2 No viability of clones and types of models

For all models where fitness affects directly the birth rate (all except Bozic), if you specify that some event (say, mutating gene A) has  $s_A \leq -1$ , if that event happens then birth rate becomes zero. This is taken to indicate that the clone is not even viable and thus disappears immediately without any chance for mutation<sup>8</sup>.

Models based on Bozic, however, have a birth rate of 1 and mutations affect the death rate. In this case, a death rate larger than birth rate, per se, does not signal

<sup>&</sup>lt;sup>8</sup>This is a shortcut that we take because we think that it is what you mean. Note, however, that technically a clone with birth rate of 0 might have a non-zero probability of mutating before becoming extinct because in the continuous time model we use mutation is not linked to reproduction. In the present code, we are not allowing for any mutation when birth rate is 0. There are other options, but none which I find really better. An alternative implementation makes a clone immediately extinct if and only if any of the  $s_i = -\infty$ . However, we still need to handle the case with  $s_i < -1$  as a special case. We either make it identical to the case with any  $s_i = -\infty$  or for any  $s_i > -\infty$  we set  $(1 + s_i) = \max(0, 1 + s_i)$  (i.e., if  $s_i < -1$  then  $(1 + s_i) = 0$ ), to avoid obtaining negative birth rates (that make no sense) and the problem of multiplying an even number of negative numbers. I think only the second would make sense as an alternative.

immediate extinction and, moreover, even for death rates that are a few times larger than birth rates, the clone could mutate before becoming extinct<sup>9</sup>.

In general, if you want to identify some mutations or some combinations of mutations as leading to immediate extinction (i.e., no viability), of the affected clone, set it to  $-\infty$  as this would work even if how birth rates of 0 are handled changes. Most examples below evaluate fitness by its effects on the birth rate. You can see one where we do it both ways in Section 3.11.1.

#### 3.3 Genes without interactions

This is a simple scenario. Each gene i has a fitness effect  $s_i$  if mutated. The  $s_i$  can come from any distribution you want. As an example let's use three genes. We know there are no order effects, but we will also see what happens if we examine genotypes as ordered.

```
ai1 <- evalAllGenotypes(allFitnessEffects(
   noIntGenes = c(0.05, -.2, .1)), order = FALSE)</pre>
```

We can easily verify the first results:

```
ai1
##
     Genotype Fitness
## 1
            1
                 1.050
## 2
            2
                 0.800
## 3
            3
                 1.100
        1, 2
## 4
                 0.840
         1, 3
## 5
                 1.155
         2, 3
## 6
                 0.880
## 7 1, 2, 3
                 0.924
```

```
all(ai1[, "Fitness"] == c( (1 + .05), (1 - .2), (1 + .1),
	(1 + .05) * (1 - .2),
	(1 + .05) * (1 + .1),
	(1 - .2) * (1 + .1),
	(1 + .05) * (1 - .2) * (1 + .1)))
## [1] TRUE
```

And we can see that considering the order of mutations (see section 3.6) makes no difference:

```
(ai2 <- evalAllGenotypes(allFitnessEffects(
    noIntGenes = c(0.05, -.2, .1)), order = TRUE,
    addwt = TRUE))
### Genotype Fitness</pre>
```

<sup>&</sup>lt;sup>9</sup>We said "a few times". For a clone of population size 1 —which is the size at which all clones start from mutation—, if death rate is, say, 90 but birth rate is 1, the probability of mutating before becoming extinct is very, very close to zero for all reasonable values of mutation rate}. How do we signal immediate extinction or no viability in this case? You can set the value of  $s = -\infty$ .

```
## 1
              WT
                   1.000
## 2
               1
                   1.050
## 3
               2
                   0.800
               3
## 4
                   1.100
          1 > 2
## 5
                   0.840
          1 > 3
## 6
                   1.155
## 7
          2 > 1
                   0.840
          2 > 3
                   0.880
## 8
          3 > 1
## 9
                   1.155
## 10
          3 > 2
                   0.880
## 11 1 > 2 > 3
                   0.924
## 12 1 > 3 > 2
                   0.924
## 13 2 > 1 > 3
                   0.924
## 14 2 > 3 > 1
                   0.924
## 15 3 > 1 > 2
                   0.924
## 16 3 > 2 > 1
                   0.924
```

(The meaning of the notation in the output table is as follows: "WT" denotes the wild-type, or non-mutated clone. The notation x > y means that a mutation in "x" happened before a mutation in "y". A genotype  $x > y \_ z$  means that a mutation in "x" happened before a mutation in "y"; there is also a mutation in "z", but that is a gene for which order does not matter).

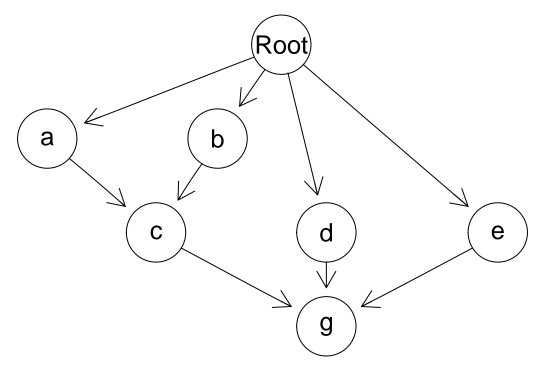
And what if I want genes without interactions but I want modules (see section 3.5)? Go to section 3.8.

# 3.4 Using DAGs: Restrictions in the order of mutations as extended posets

#### 3.4.1 AND, OR, XOR relationships

The literature on Oncogenetic trees, CBNs, etc, has used graphs as a way of showing the restrictions in the order in which mutations can accumulate. The meaning of "convergent arrows" in these graphs, however, differs. In Figure 1 of Korsunsky et al. (2014) we are shown a simple diagram that illustrates the three basic different meanings of convergent arrows using two parental nodes. We will illustrate it here with three. Suppose we focus on node "g" in the following figure (we will create it shortly)

```
data(examplesFitnessEffects)
plot(examplesFitnessEffects[["cbn1"]])
```



- In relationships of the type used in Conjunctive Bayesian Networks (CBN) (e.g., Gerstung et al. 2009), we are modeling an AND relationship, also called CMPN by Korsunsky et al. (2014) or monotone relationship by Farahani and Lagergren (2013). If the relationship in the graph is fully respected, then "g" will only appear if all of "c", "d", and "e" are already mutated.
- Semimonotone relationships sensu Farahani and Lagergren (2013) or **DMPN** sensu Korsunsky et al. (2014) are **OR** relationships: "g" will appear if one or more of "c", "d", or "e" are already mutated.
- XMPN relationships (Korsunsky et al. 2014) are XOR relationships: "g" will be present only if exactly one of "c", "d", or "e" is present.

Note that Oncogenetic trees (Desper et al. 1999; Szabo and Boucher 2008) need not deal with the above distinctions, since the DAGs are trees: no node has more than one incoming connection or more than one parent<sup>10</sup>.

To have a flexible way of specifying all of these restrictions, we will want to be able to say what kind of dependency each child node has on its parents.

## 3.4.2 Fitness effects

Those DAGs specify dependencies and, as explained in Diaz-Uriarte (2015), it is simple to map them to a simple evolutionary model: any set of mutations that does not conform to the restrictions encoded in the graph will have a fitness of 0. However, we might not want to require absolute compliance with the DAG. This means we might want to allow deviations from the DAG with a corresponding penalization

 $<sup>^{10}{</sup>m OTs}$  and CBNs have some other technical differences about the underlying model they assume, such as the exponential waiting time in CBNs. We will not discuss them here.

that is, however, not identical to setting fitness to 0 (again, see Diaz-Uriarte 2015). This we can do by being explicit about the fitness effects of the deviations from the restrictions encoded in the DAG. We will use below a column of s for the fitness effect when the restrictions are satisfied and a column of sh when they are not. (See also 3.2 for the details about the meaning of the fitness effects).

That way of specifying fitness effects makes it also trivial to use the model in Hjelm, Höglund, and Lagergren (2006) where all mutations might be allowed to occur, but the presence of some mutations increases the probability of occurrence of other mutations. For example, the values of sh could be all small positive ones (or for mildly deleterious effects, small negative numbers), while the values of s are much larger positive numbers.

#### 3.4.3 Extended posets

In version 1 of this package we used posets in the sense of Beerenwinkel, Eriksson, and Sturmfels (2007) and Gerstung et al. (2009), as explained in section 15.1 and in the help for poset. Here, we continue using two columns, that specify parents and children, but we add columns for the specific values of fitness effects (both s and sh—i.e., fitness effects for what happens when restrictions are and are not satisfied) and for the type of dependency as explained in section 3.4.1.

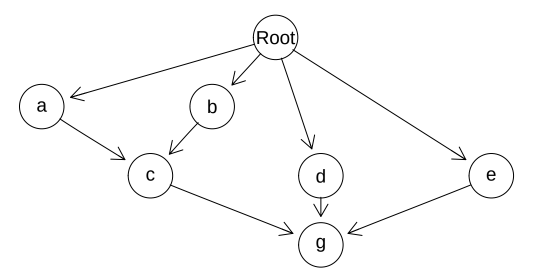
We can now illustrate the specification of different fitness effects using DAGs.

## 3.4.4 DAGs: A first conjunction (AND) example

(We skip one letter, just to show that names need not be consecutive or have any particular order.)

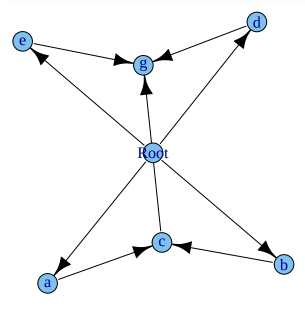
We can get a graphical representation using the default "graphNEL"

```
plot(cbn1)
```



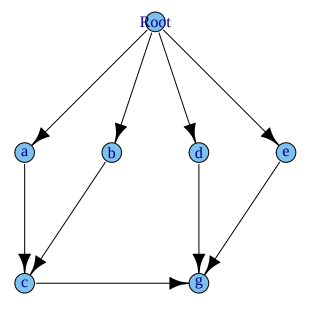
or one using "igraph":

## plot(cbn1, "igraph")



Since we have a parent and children, the reingold.tilford layout is probably the best here, so you might want to use that:

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



And what is the fitness of all genotypes?

```
gfs <- evalAllGenotypes(cbn1, order = FALSE, addwt = TRUE)
gfs[1:15,]
##
       Genotype Fitness
              WT
                     1.00
## 1
## 2
                     1.10
               \boldsymbol{a}
## 3
                     1.10
               b
                     0.10
## 4
               С
                     1.10
## 5
               d
## 6
                     1.10
               е
## 7
                     0.10
               g
## 8
           a,
               b
                     1.21
## 9
                     0.11
               С
            а,
## 10
                     1.21
               d
## 11
                     1.21
               е
## 12
                     0.11
              g
                     0.11
## 13
            b, c
## 14
            b, d
                     1.21
## 15
           b, e
                     1.21
```

You can verify that for each genotype, if a mutation is present without all of its dependencies present, you get a (1-0.9) multiplier, and you get a (1+0.1) multiplier for all the rest with its direct parents satisfied. For example, genotypes "a", or "b", or "d", or "e" have fitness (1+0.1), genotype "a, b, c" has fitness  $(1+0.1)^3$ , but genotype "a, c" has fitness (1+0.1)(1-0.9) = 0.11.

## 3.4.5 DAGs: A second conjunction example

Let's try a first attempt at a somewhat more complex example, where the fitness consequences of different genes differ.

If you try this, you'll get an error. There is an error because the "sh" varies within a child, and we do not allow that for a poset-type specification, as it is ambiguous. If you need arbitrary fitness values for arbitrary combinations of genotypes, you can specify them using epistatic effects as in section 3.7 and order effects as in section 3.6.

Why do we need to specify as many "s" and "sh" as there are rows (or a single one, that gets expanded to those many) when the "s" and "sh" are properties of the child node, not of the edges? Because, for ease, we use a data.frame.

We fix the error in our specification. Notice that the "sh" is not set to -1 in these examples. If you want strict compliance with the poset restrictions, you should set sh=-1 or, better yet,  $sh=-\infty$  (see section 3.2.2), but having an sh>-1 will lead to fitnesses that are >0 and, thus, is a way of modeling small deviations from the poset (see discussion in Diaz-Uriarte 2015).

Note that for those nodes that depend only on "Root" the type of dependency is irrelevant.

We could get graphical representations but the figures would be the same as in the example in section 3.4.4, since the structure has not changed, only the numeric values.

What is the fitness of all possible genotypes? Here, order of events *per se* does not matter, beyond that considered in the poset. In other words, the fitness of genotype "a, b, c" is the same no matter how we got to "a, b, c". What matters is whether or not the genes on which each of "a", "b", and "c" depend are present or not (I only show the first 10 genotypes)

```
gcbn2 <- evalAllGenotypes(cbn2, order = FALSE)
gcbn2[1:10, ]
## Genotype Fitness</pre>
```

```
## 1
                  1.010
## 2
                  1.020
## 3
                  0.100
             С
## 4
             d
                  1.030
## 5
                  1.040
             е
## 6
                  0.050
             g
## 7
                  1.030
          a, b
                  0.101
## 8
          a, c
## 9
          a, d
                  1.040
## 10
          a, e
                  1.050
```

Of course, if we were to look at genotypes but taking into account order of occurrence of mutations, we would see no differences

```
gcbn2o <- evalAllGenotypes(cbn2, order = TRUE, max = 1956)
gcbn2o[1:10, ]
##
      Genotype Fitness
                  1.010
## 1
             \boldsymbol{a}
## 2
                 1.020
             b
## 3
                 0.100
             C
## 4
             d
                 1.030
                 1.040
## 5
             e
## 6
                 0.050
             g
## 7
         a > b
                 1.030
## 8
         a > c
                 0.101
## 9
         a > d
                 1.040
## 10
         a > e 1.050
```

(The max = 1956 is there so that we show all the genotypes, even if they are more than 256, the default.)

You can check the output and verify things are as they should. For instance:

```
all.equal(
    gcbn2[c(1:21, 22, 28, 41, 44, 56, 63), "Fitness"],
    c(1.01, 1.02, 0.1, 1.03, 1.04, 0.05,
        1.01 * c(1.02, 0.1, 1.03, 1.04, 0.05),
        1.02 * c(0.10, 1.03, 1.04, 0.05),
        0.1 * c(1.03, 1.04, 0.05),
        1.03 * c(1.04, 0.05),
        1.04 * 0.05,
        1.01 * 1.02 * 1.1,
        1.01 * 0.1 * 0.05,
        1.03 * 1.04 * 0.05,
        1.03 * 1.04 * 0.05,
        1.01 * 1.02 * 1.1 * 0.05,
        1.03 * 1.04 * 1.2 * 0.1, ## notice this
        1.01 * 1.02 * 1.03 * 1.04 * 1.1 * 1.2
        ))
```

## ## [1] TRUE

A particular one that is important to understand is genotype with mutated genes "c, d, e, g":

```
gcbn2[56,]

## Genotype Fitness

## 56 c, d, e, g 0.1285

all.equal(gcbn2[56, "Fitness"], 1.03 * 1.04 * 1.2 * 0.10)

## [1] TRUE
```

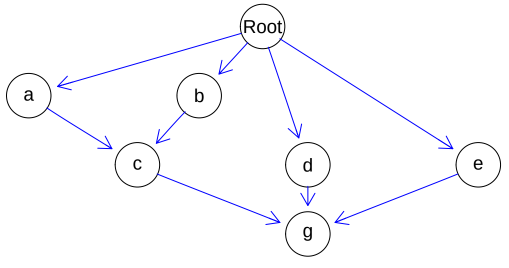
where "g" is taken as if its dependencies are satisfied (as "c", "d", and "e" are present) even when the dependencies of "c" are not satisfied (and that is why the term for "c" is 0.9).

## 3.4.6 DAGs: A semimonotone or "OR" example

We will reuse the above example, changing the type of relationship:

It looks like this (where edges are shown in blue to denote the semimonotone relationship):

#### plot(smn1)



```
gsmn1 <- evalAllGenotypes(smn1, order = FALSE)</pre>
```

Having just one parental dependency satisfied is now enough, in contrast to what

happened before. For instance:

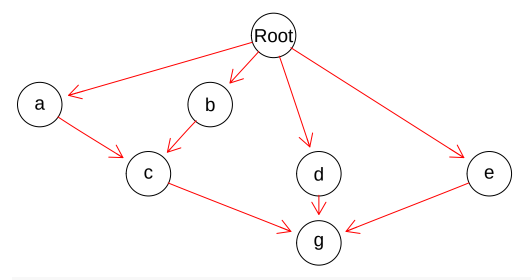
```
gcbn2[c(8, 12, 22),]
##
     Genotype Fitness
## 8
         a, c
                0.101
         b, c
## 12
                0.102
## 22 a, b, c 1.133
gsmn1[c(8, 12, 22),]
    Genotype Fitness
## 8
               1.111
         a, c
## 12
        b, c 1.122
## 22 a, b, c 1.133
gcbn2[c(20:21, 28),]
     Genotype Fitness
         d, g 0.05150
## 20
## 21
         e, g 0.05200
## 28 a, c, g 0.00505
gsmn1[c(20:21, 28),]
##
    Genotype Fitness
## 20
         d, g 1.236
## 21
         e, g 1.248
## 28 a, c, g 1.333
```

## 3.4.7 An "XMPN" or "XOR" example

Again, we reuse the example above, changing the type of relationship:

It looks like this (edges in red to denote the "XOR" relationship):

```
plot(xor1)
```



```
gxor1 <- evalAllGenotypes(xor1, order = FALSE)</pre>
```

Whenever "c" is present with both "a" and "b", the fitness component for "c" will be (1-0.1). Similarly for "g" (if more than one of "d", "e", or "c" is present, it will show as (1-0.05)). For example:

```
gxor1[c(22, 41), ]

## Genotype Fitness

## 22 a, b, c 0.10302

## 41 d, e, g 0.05356

c(1.01 * 1.02 * 0.1, 1.03 * 1.04 * 0.05)

## [1] 0.10302 0.05356
```

However, having just both "a" and "b" is identical to the case with CBN and the monotone relationship (see sections 3.4.5 and 3.4.6). If you want the joint presence of "a" and "b" to result in different fitness than the product of the individual terms, without considering the presence of "c", you can specify that using general epistatic effects (section 3.7).

We also see a very different pattern compared to CBN (section 3.4.5) here:

```
gxor1[28, ]
## Genotype Fitness
## 28 a, c, g 1.333
1.01 * 1.1 * 1.2
## [1] 1.333
```

as exactly one of the dependencies for both "c" and "g" are satisfied.

But

```
gxor1[44,]

## Genotype Fitness

## 44 a, b, c, g 0.1236

1.01 * 1.02 * 0.1 * 1.2
```

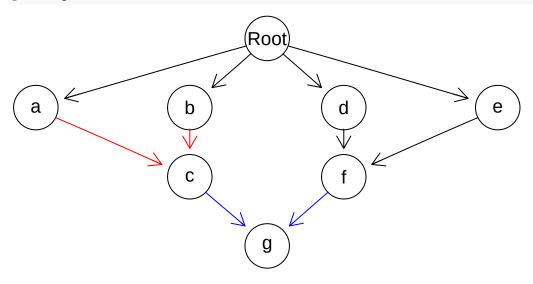
## ## [1] 0.1236

is the result of a 0.1 for "c" (and a 1.2 for "g" that has exactly one of its dependencies satisfied).

## 3.4.8 Posets: the three types of relationships

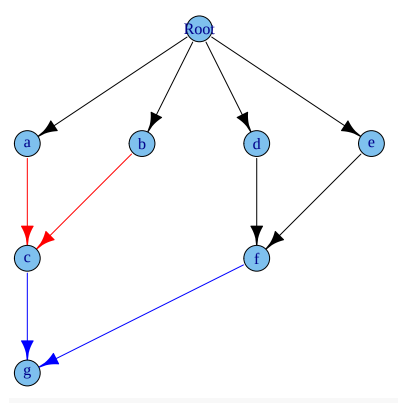
This is how it looks like:

## plot(fp3)



We can also use "igraph":

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



gfp3 <- evalAllGenotypes(fp3, order = FALSE)</pre>

Let's look at a few:

```
gfp3[c(9, 24, 29, 59, 60, 66, 119, 120, 126, 127), ]
##
                  Genotype Fitness
## 9
                      a, c 1.111000
                      d, f 0.051500
## 24
## 29
                   a, b, c 0.103020
## 59
                   c, f, g 0.006500
## 60
                   d, e, f 1.285440
## 66
                a, b, c, f 0.005151
             c, d, e, f, g 0.167107
## 119
## 120
          a, b, c, d, e, f 0.132426
## 126
          b, c, d, e, f, g 1.874943
## 127 a, b, c, d, e, f, g 0.172154
c(1.01 * 1.1, 1.03 * .05, 1.01 * 1.02 * 0.1, 0.1 * 0.05 * 1.3,
 1.03 * 1.04 * 1.2, 1.01 * 1.02 * 0.1 * 0.05,
 0.1 * 1.03 * 1.04 * 1.2 * 1.3,
 1.01 * 1.02 * 0.1 * 1.03 * 1.04 * 1.2,
 1.02 * 1.1 * 1.03 * 1.04 * 1.2 * 1.3,
 1.01 * 1.02 * 1.03 * 1.04 * 0.1 * 1.2 * 1.3)
  [1] 1.111000 0.051500 0.103020 0.006500 1.285440 0.005151 0.167107
   [8] 0.132426 1.874943 0.172154
```

As before, looking at the order of mutations makes no difference (look at the test

directory to see a test that verifies this assertion).

#### 3.5 Modules

As already mentioned, we can think of all the effects of fitness in terms not of individual genes but, rather, modules. This idea is discussed in, for example, Raphael and Vandin (2015), Gerstung et al. (2011): the restrictions encoded in, say, the DAGs can be considered to apply not to genes, but to modules, where each module is a set of genes (and the intersection between modules is the empty set). Modules, then, play the role of a "union operation" over sets of genes. Of course, if we can use modules for the restrictions in the DAGs we should also be able to use them for epistasis and order effects, as we will see later (e.g., 3.6.2).

## 3.5.1 What does a module provide

Modules can provide very compact ways of specifying relationships when you want to, well, model the existence of modules. For simplicity suppose there is a module, "A", made of genes "a1" and "a2", and a module "B", made of a single gene "b1". Module "B" can mutate if module "A" is mutated, but mutating both "a1" and "a2" provides no additional fitness advantage compared to mutating only a single one of them. We can specify this as:

```
s < -0.2
sboth < (1/(1 + s)) - 1
m0 <- allFitnessEffects(data.frame(</pre>
    parent = c("Root", "Root", "a1", "a2"),
    child = c("a1", "a2", "b", "b"),
    s = s,
    sh = -1,
    typeDep = "OR"),
                         epistasis = c("a1:a2" = sboth))
evalAllGenotypes(m0, order = FALSE, addwt = TRUE)
##
      Genotype Fitness
             WT
## 1
                   1.00
                   1.20
## 2
             a1
## 3
             a2
                   1.20
## 4
              b
                   0.00
        a1, a2
                   1.20
## 5
         a1, b
## 6
                   1.44
## 7
         a2, b
                   1.44
## 8 a1, a2, b
                   1.44
```

Note that we need to add an epistasis term, with value "sboth" to capture the idea of "mutating both"a1" and "a2" provides no additional fitness advantage compared to mutating only a single one of them"; see details in section 3.7.

Now, specify it using modules:

```
s < -0.2
m1 <- allFitnessEffects(data.frame(</pre>
    parent = c("Root", "A"),
    child = c("A", "B"),
    s = s,
    sh = -1,
    typeDep = "OR"),
                         geneToModule = c("Root" = "Root",
                                            "A" = "a1, a2",
                                            "B" = "b1"))
evalAllGenotypes(m1, order = FALSE, addwt = TRUE)
       Genotype Fitness
##
## 1
              WT
                     1.00
## 2
                     1.20
              a1
## 3
              a2
                     1.20
## 4
              b1
                     0.00
## 5
         a1, a2
                     1.20
## 6
         a1, b1
                     1.44
## 7
         a2, b1
                     1.44
## 8 a1, a2, b1
                     1.44
```

This captures the ideas directly. The typing savings here are small, but they can be large with modules with many genes.

#### 3.5.2 Specifying modules

How do you specify modules? The general procedure is simple: you pass a vector that makes explicit the mapping from modules to sets of genes. We just saw an example. There are several additional examples such as 3.5.3, 3.6.2, 3.7.4.

It is important to note that, once you specify modules, we expect all of the relationships (except those that involve the non interacting genes) to be specified as modules. Thus, all elements of the epistasis, posets (the DAGs) and order effects components should be specified in terms of modules. But you can, of course, specify a module as containing a single gene (and a single gene with the same name as the module).

What about the "Root" node? If you use a "restriction table", that restriction table (that DAG) must have a node named "Root" and in the mapping of genes to module there **must** be a first entry that has a module and gene named "Root", as we saw above with <code>geneToModule = c("Root" = "Root", ....</code> We force you to do this to be explicit about the "Root" node. This is not needed (thought it does not hurt) with other fitness specifications. For instance, if we have a model with two modules, one of them with two genes (see details in section 3.8) we do not need to pass a "Root" as in

```
fnme <- allFitnessEffects(epistasis = c("A" = 0.1, "B" = 0.2), geneToModule = c("A" = "a1, a2",
```

```
"B" = "b1"))
evalAllGenotypes(fnme, order = FALSE, addwt = TRUE)
##
       Genotype Fitness
## 1
              WT
                    1.00
                    1.10
## 2
              a1
## 3
              a2
                    1.10
## 4
              b1
                    1.20
## 5
         a1, a2
                    1.10
## 6
         a1, b1
                    1.32
## 7
         a2, b1
                    1.32
## 8 a1, a2, b1
                    1.32
```

but it is also OK to have a "Root" in the geneToModule:

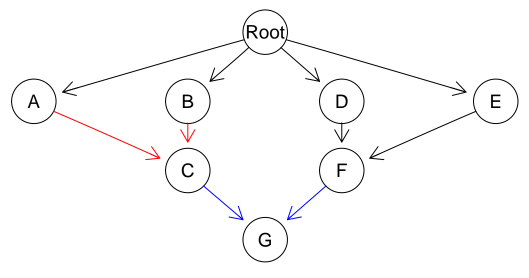
```
fnme2 <- allFitnessEffects(epistasis = c("A" = 0.1,</pre>
                                           "B" = 0.2),
                           geneToModule = c(
                                "Root" = "Root",
                                "A" = "a1, a2",
                                "B" = "b1"))
evalAllGenotypes(fnme, order = FALSE, addwt = TRUE)
##
       Genotype Fitness
## 1
              WT
                    1.00
## 2
              a1
                    1.10
## 3
              a2
                    1.10
## 4
                    1.20
              b1
## 5
                    1.10
         a1, a2
         a1. b1
## 6
                    1.32
## 7
         a2, b1
                    1.32
## 8 a1, a2, b1
                    1.32
```

## 3.5.3 Modules and posets again: the three types of relationships and modules

We use the same specification of poset, but add modules. To keep it manageable, we only add a few genes for some modules, and have some modules with a single gene. Beware that the number of genotypes is starting to grow quite fast, though. We capitalize to differentiate modules (capital letters) from genes (lowercase with a number), but this is not needed.

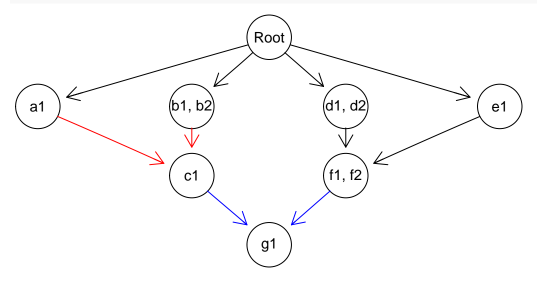
By default, plotting shows the modules:

## plot(fp4m)

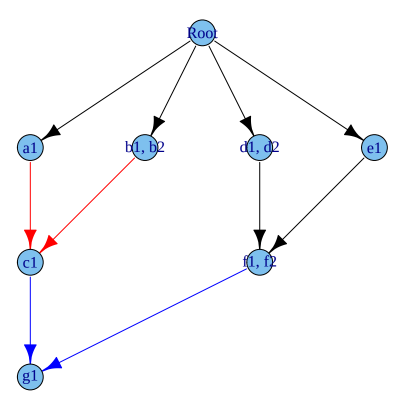


but we can show the gene names instead of the module names:

## plot(fp4m, expandModules = TRUE)



or



We obtain the fitness of all genotypes in the usual way:

```
gfp4 <- evalAllGenotypes(fp4m, order = FALSE, max = 1024)</pre>
```

Let's look at a few of those:

```
gfp4[c(12, 20, 21, 40, 41, 46, 50, 55, 64, 92,
       155, 157, 163, 372, 632, 828), ]
##
                      Genotype Fitness
## 12
                        a1, b2
                               1.0302
## 20
                        b1, b2
                                1.0200
## 21
                        b1, c1
                                1.1220
                        c1, g1
## 40
                                0.1300
## 41
                        d1, d2
                                1.0300
## 46
                        d2, e1
                                1.0712
## 50
                        e1, f1
                                0.0520
                                0.0650
## 55
                        f2, g1
                    a1, b2, c1
## 64
                                0.1030
                    b1, b2, c1
## 92
                                1.1220
## 155
                    c1, f2, g1
                                0.0065
                    d1, d2, f1
## 157
                                0.0515
## 163
                    d1, f1, f2
                               0.0515
## 372
               d1, d2, e1, f2
                                1.2854
## 632
           d1, d2, e1, f1, f2
                               1.2854
## 828 b2, c1, d1, e1, f2, g1
                                1.8749
c(1.01 * 1.02, 1.02, 1.02 * 1.1, 0.1 * 1.3, 1.03,
```

```
1.03 * 1.04, 1.04 * 0.05, 0.05 * 1.3,

1.01 * 1.02 * 0.1, 1.02 * 1.1, 0.1 * 0.05 * 1.3,

1.03 * 0.05, 1.03 * 0.05, 1.03 * 1.04 * 1.2, 1.03 * 1.04 * 1.2,

1.02 * 1.1 * 1.03 * 1.04 * 1.2 * 1.3)

## [1] 1.0302 1.0200 1.1220 0.1300 1.0300 1.0712 0.0520 0.0650 0.1030

## [10] 1.1220 0.0065 0.0515 0.0515 1.2854 1.2854 1.8749
```

## 3.6 Order effects

As explained in the introduction (section 1), by order effects we mean a phenomenon such as the one shown empirically by Ortmann et al. (2015): the fitness of a double mutant "A", "B" is different depending on whether "A" was acquired before "B" or "B" before "A". This, of course, can be generalized to more than two genes.

Note that order effects are different from the restrictions in the order of accumulation of mutations discussed in section 3.4. With restrictions in the order of accumulation of mutations we might say that acquiring "B" depends or is facilitated by having "A" mutated (and, unless we allowed for multiple mutations, having "A" mutated means having "A" mutated before "B"). However, once you have the genotype "A, B", its fitness does not depend on the order in which "A" and "B" appeared.

## 3.6.1 Order effects: three-gene orders

Consider this case, where three specific three-gene orders and two two-gene orders (one of them a subset of one of the three) lead to different fitness compared to the wild-type. We add also modules, to show its usage (but just limit ourselves to using one gene per module here).

Order effects are specified using a x > y, which means that that order effect is satisfied when module x is mutated before module y.

```
o3 <- allFitnessEffects(orderEffects = c(
                                   "F > D > M" = -0.3,
                                   "D > F > M" = 0.4,
                                   "D > M > F" = 0.2.
                                   "D > M"
                                                  = 0.1,
                                                  = 0.5),
                                   "M > D"
                              geneToModule =
                                   c("M" = "m",
                                     ^{\prime\prime}F^{\prime\prime} = ^{\prime\prime}f^{\prime\prime}
                                     "D" = "d") )
(ag <- evalAllGenotypes(o3, addwt = TRUE, order = TRUE))
         Genotype Fitness
## 1
                WT
                        1.00
## 2
                 d
                        1.00
```

```
## 3
                    1.00
## 4
                    1.00
              m
## 5
          d > f
                    1.00
## 6
          d > m
                    1.10
## 7
          f > d
                    1.00
## 8
                    1.00
          f > m
                    1.50
## 9
          m > d
                    1.00
## 10
          m > f
## 11 d > f > m
                    1.54
## 12 d > m > f
                    1.32
## 13 f > d > m
                    0.77
## 14 f > m > d
                    1.50
## 15 m > d > f
                    1.50
## 16 m > f > d
                    1.50
```

(The meaning of the notation in the output table is as follows: "WT" denotes the wild-type, or non-mutated clone. The notation x > y means that a mutation in "x" happened before a mutation in "y". A genotype  $x > y \_ z$  means that a mutation in "x" happened before a mutation in "y"; there is also a mutation in "z", but that is a gene for which order does not matter).

The values for the first nine genotypes come directly from the fitness specifications. The 10th genotype matches D > F > M (= (1 + 0.4)) but also D > M ((1 + 0.1)). The 11th matches D > M > F and D > M. The 12th matches F > D > M but also D > M. Etc.

#### 3.6.2 Order effects and modules with multiple genes

Consider the following case:

There are four genes, d1, d2, f1, f2, where each d belongs to module D and each f belongs to module F.

What to expect for cases such as d1 > f1 or f1 > d1 is clear, as shown in

```
ag[5:16,]

## Genotype Fitness

## 5 d1 > d2 1.0

## 6 d1 > f1 1.4

## 7 d1 > f2 1.4
```

```
## 8
       d2 > d1
                    1.0
## 9
       d2 > f1
                    1.4
## 10
       d2 > f2
                    1.4
## 11
       f1 > d1
                    0.7
## 12
       f1 > d2
                    0.7
## 13
       f1 > f2
                    1.0
## 14
       f2 > d1
                    0.7
## 15
                    0.7
       f2 > d2
## 16
       f2 > f1
                    1.0
```

Likewise, cases such as d1 > d2 > f1 or f2 > f1 > d1 are clear, because in terms of modules they map to \$D > F\$ or F > D: the observed order of mutation d1 > d2 > f1 means that module D was mutated first and module F was mutated second. Similar for d1 > f2 or f1 > d1 > d2: those map to D > F and F > D. We can see the fitness of those four case in:

and they correspond to the values of those order effects, where F > D = (1 - 0.3) and D > F = (1 + 0.4):

```
ag[c(17, 39, 19, 29), "Fitness"] == c(1.4, 0.7, 1.4, 0.7)
## [1] TRUE TRUE TRUE
```

What if we match several patterns? For example, d1 > f1 > d2 > f2 and d1 > f1 > f2 > d2? The first maps to D > F > D > F and the second to D > F > D. But since we are concerned with which one happened first and which happened second we should expect those two to correspond to the same fitness, that of pattern D > F, as is the case:

More generally, that applies to all the patterns that start with one of the "d" genes:

```
all(ag[41:52, "Fitness"] == 1.4)
## [1] TRUE
```

Similar arguments apply to the opposite pattern, F > D, which apply to all the possible gene mutation orders that start with one of the "f" genes. For example:

```
all(ag[53:64, "Fitness"] == 0.7)
## [1] TRUE
```

#### 3.6.3 Order and modules with 325 genotypes

We can of course have more than two genes per module. This just repeats the above, with five genes (there are 325 genotypes, and that is why we pass the "max" argument to evalAllGenotypes, to allow for more than the default 256).

We can verify that any combination that starts with a "d" gene and then contains at least one "f" gene will have a fitness of 1 + 0.4. And any combination that starts with an "f" gene and contains at least one "d" genes will have a fitness of 1 - 0.3. All other genotypes have a fitness of 1:

```
all(ag2[grep("^d.*f.*", ag2[, 1]), "Fitness"] == 1.4)
## [1] TRUE
all(ag2[grep("^f.*d.*", ag2[, 1]), "Fitness"] == 0.7)
## [1] TRUE
oe <- c(grep("^f.*d.*", ag2[, 1]), grep("^d.*f.*", ag2[, 1]))
all(ag2[-oe, "Fitness"] == 1)
## [1] TRUE</pre>
```

## 3.6.4 Order effects and genes without interactions

We will now look at both order effects and interactions. To make things more interesting, we name genes so that the ordered names do split nicely between those with and those without order effects (this, thus, also serves as a test of messy orders of names).

```
foi1 <- allFitnessEffects(
    orderEffects = c("D>B" = -0.2, "B > D" = 0.3),
    noIntGenes = c("A" = 0.05, "C" = -.2, "E" = .1))
```

You can get a verbose view of what the gene names and modules are (and their automatically created numeric codes) by:

```
foi1[c("geneModule", "long.geneNoInt")]
## $geneModule
## Gene Module GeneNumID ModuleNumID
## 1 Root Root 0 0
## 2 B B 1 1
```

```
## 3 D D 2 2
##

## $long.geneNoInt

## Gene GeneNumID s

## A A 3 0.05

## C C 4 -0.20

## E E 5 0.10
```

We can get the fitness of all genotypes (we set max = 325 because that is the number of possible genotypes):

```
agoi1 <- evalAllGenotypes(foi1, max = 325, order = TRUE)
head(agoi1)
##
      Genotype Fitness
## 1
              В
                    1.00
## 2
              D
                    1.00
## 3
              \boldsymbol{A}
                    1.05
## 4
              C
                    0.80
## 5
              \boldsymbol{E}
                    1.10
## 6
         B > D
                    1.30
```

Now:

```
rn <- 1:nrow(agoi1)
names(rn) <- agoi1[, 1]

agoi1[rn[LETTERS[1:5]], "Fitness"] == c(1.05, 1, 0.8, 1, 1.1)
## [1] TRUE TRUE TRUE TRUE</pre>
```

According to the fitness effects we have specified, we also know that any genotype with only two mutations, one of which is either "A", "C" "E" and the other is "B" or "D" will have the fitness corresponding to "A", "C" or "E", respectively:

```
agoi1[grep("^A > [BD]$", names(rn)), "Fitness"] == 1.05
## [1] TRUE TRUE
agoi1[grep("^C > [BD]$", names(rn)), "Fitness"] == 0.8
## [1] TRUE TRUE
agoi1[grep("^E > [BD]$", names(rn)), "Fitness"] == 1.1
## [1] TRUE TRUE
agoi1[grep("^[BD] > A$", names(rn)), "Fitness"] == 1.05
## [1] TRUE TRUE
agoi1[grep("^[BD] > C$", names(rn)), "Fitness"] == 0.8
## [1] TRUE TRUE
agoi1[grep("^[BD] > E$", names(rn)), "Fitness"] == 1.1
## [1] TRUE TRUE
```

We will not be playing many additional games with regular expressions, but let us check those that start with "D" and have all the other mutations, which occupy rows 230 to 253; fitness should be equal (within numerical error, because of floating

point arithmetic) to the order effect of "D" before "B" times the other effects (1-0.3)\*1.05\*0.8\*1.1=0.7392

and that will also be the value of any genotype with the five mutations where "D" comes before "B" such as those in rows 260 to 265, 277, or 322 and 323, but it will be equal to (1+0.3)\*1.05\*0.8\*1.1 = 1.2012 in those where "B" comes before "D". Analogous arguments apply to four, three, and two mutation genotypes.

## 3.7 Epistasis

## 3.7.1 Epistasis: two alternative specifications

We want the following mapping of genotypes to fitness:

Ā	В	Fitness
wt	wt	1
wt	M	$1+s_b$
Μ	wt	$1+s_a$
Μ	M	$1 + s_{ab}$

Suppose that the actual numerical values are  $s_a = 0.2, s_b = 0.3, s_{ab} = 0.7$ .

We specify the above as follows:

```
sa <- 0.2
sb <- 0.3
sab <- 0.7
e2 <- allFitnessEffects(epistasis =</pre>
                               c("A: -B" = sa,
                                  "-A:B" = sb,
                                  "A : B" = sab)
evalAllGenotypes(e2, order = FALSE, addwt = TRUE)
##
     Genotype Fitness
            WT
## 1
                     1.0
## 2
                     1.2
             \boldsymbol{A}
## 3
             В
                     1.3
                    1.7
          A, B
```

That uses the "-" specification, so we explicitly exclude some patterns: with "A:-B" we say "A when there is no B".

But we can also use a specification where we do not use the "-". That requires a different numerical value of the interaction, because now, as we are rewriting the

interaction term as genotype "A is mutant, B is mutant" the double mutant will incorporate the effects of "A mutant", "B mutant" and "both A and B mutants". We can define a new  $s_2$  that satisfies  $(1+s_{ab})=(1+s_a)(1+s_b)(1+s_2)$  so  $(1+s_2)=(1+s_{ab})/((1+s_a)(1+s_b))$  and therefore specify as:

```
s2 \leftarrow ((1 + sab)/((1 + sa) * (1 + sb))) - 1
e3 <- allFitnessEffects(epistasis =</pre>
                               c("A" = sa,
                                 "B" = sb,
                                 "A : B" = s2)
evalAllGenotypes(e3, order = FALSE, addwt = TRUE)
     Genotype Fitness
## 1
            WT
                    1.0
## 2
             Α
                    1.2
## 3
             B
                    1.3
                    1.7
## 4
          A, B
```

Note that this is the way you would specify effects with FFPopsim (Zanini and Neher 2012). Whether this specification or the previous one with "-" is simpler will depend on the model. For synthetic mortality and viability, I think the one using "-" is simpler to map genotype tables to fitness effects. See also section 3.7.2 and 3.7.3 and the example in section 5.4.2.

Finally, note that we can also specify some of these effects by combining the graph and the epistasis, as shown in section 5.2.1 or 5.4.2.

#### 3.7.2 Epistasis with three genes and two alternative specifications

Suppose we have

A	В	С	Fitness
M	wt	wt	$1+s_a$
wt	M	wt	$1 + s_b$
wt	wt	M	$1+s_c$
Μ	M	wt	$1 + s_{ab}$
wt	Μ	M	$1+s_{bc}$
Μ	wt	M	$(1+s_a)(1+s_c)$
Μ	Μ	M	$1 + s_{abc}$

where missing rows have a fitness of 1 (they have been deleted for conciseness). Note that the mutant for exactly A and C has a fitness that is the product of the individual terms (so there is no epistasis in that case).

```
sa <- 0.1
sb <- 0.15
sc <- 0.2
```

```
sab <- 0.3
sbc < -0.25
sabc <- 0.4
sac \leftarrow (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)
##
     Genotype Fitness
## 1
             \boldsymbol{A}
                   1.10
## 2
             B
                   1.15
## 3
             C
                   1.20
## 4
         A, B
                  1.30
## 5
         A, C
                   1.32
                   0.75
## 6
         B, C
## 7 A, B, C
                   1.40
```

We needed to pass the  $s_{ac}$  coefficient explicitly, even if it that term was just the product. We can try to avoid using the "-", however (but we will need to do other calculations). For simplicity, I use capital "S" in what follows where the letters differ from the previous specification:

```
"B:C" = Sbc,
                                ## "A:C" = sac, ## not needed now
                                "A : B : C" = Sabc)
evalAllGenotypes(E3B, order = FALSE, addwt = FALSE)
     Genotype Fitness
            Α
## 1
                  1.10
## 2
            B
                  1.15
            C
## 3
                 1.20
## 4
         A, B
                 1.30
## 5
         A, C
                 1.32
         B, C
                  0.75
## 6
                 1.40
## 7 A, B, C
```

The above two are, of course, identical:

```
all(evalAllGenotypes(E3A, order = FALSE, addwt = FALSE) ==
    evalAllGenotypes(E3B, order = FALSE, addwt = FALSE))
## [1] TRUE
```

We avoid specifying the "A:C", as it just follows from the individual "A" and "C" terms, but given a specified genotype table, we need to do a little bit of addition and multiplication to get the coefficients.

## 3.7.3 Why can we specify some effects with a "-"?

Let's suppose we want to specify the synthetic viability example seen before:

A	В	Fitness
wt	wt	1
wt	${\bf M}$	0
M	wt	0
Μ	Μ	(1 + s)

where "wt" denotes wild type and "M" denotes mutant.

If you want to directly map the above table to the fitness table for the program, to specify the genotype "A is wt, B is a mutant" you can specify it as "-A,B", not just as "B". Why? Because just the presence of a "B" is also compatible with genotype "A is mutant and B is mutant". If you use "-" you are explicitly saying what should not be there so that "-A,B" is NOT compatible with "A, B". Otherwise, you need to carefully add coefficients. Depending on what you are trying to model, different specifications might be simpler. See the examples in section 3.7.1 and 3.7.2. You have both options.

# 3.7.4 Epistasis: modules

There is nothing conceptually new, but we will show an example here:

```
sa <- 0.2
sb <- 0.3
sab <- 0.7
em <- allFitnessEffects(epistasis =</pre>
                             c("A: -B" = sa,
                               "-A:B" = sb,
                               "A : B" = sab),
                         geneToModule = c("A" = "a1, a2",
                                           "B" = "b1, b2")
evalAllGenotypes(em, order = FALSE, addwt = TRUE)
##
             Genotype Fitness
## 1
                   WT
                           1.0
## 2
                   a1
                          1.2
## 3
                   a2
                          1.2
                          1.3
## 4
                   b1
                          1.3
## 5
                   b2
## 6
               a1, a2
                          1.2
## 7
               a1, b1
                          1.7
## 8
               a1, b2
                          1.7
## 9
               a2, b1
                          1.7
## 10
               a2, b2
                          1.7
               b1, b2
                          1.3
## 11
## 12
          a1, a2, b1
                          1.7
## 13
          a1, a2, b2
                          1.7
## 14
          a1, b1, b2
                          1.7
## 15
          a2, b1, b2
                          1.7
## 16 a1, a2, b1, b2
                          1.7
```

Of course, we can do the same thing without using the "-", as in section 3.7.1:

```
## 2
                           1.2
                    a1
## 3
                    a2
                           1.2
## 4
                    b1
                           1.3
## 5
                    b2
                           1.3
## 6
                           1.2
               a1, a2
## 7
               a1, b1
                           1.7
## 8
               a1, b2
                           1.7
                           1.7
## 9
               a2, b1
## 10
               a2, b2
                           1.7
## 11
               b1. b2
                           1.3
                           1.7
## 12
           a1, a2, b1
           a1, a2, b2
                           1.7
## 13
                           1.7
## 14
           a1, b1, b2
           a2, b1, b2
                           1.7
## 15
## 16 a1, a2, b1, b2
                           1.7
```

# 3.8 I do not want epistasis, but I want modules!

Sometimes you might want something like having several modules, say "A" and "B", each with a number of genes, but with "A" and "B" showing no interaction.

It is a terminological issue whether we should allow noIntGenes (no interaction genes), as explained in section 3.3 to actually be modules. The reasoning for not allowing them is that the situation depicted above (several genes in module A, for example) actually is one of interaction: the members of "A" are combined using an "OR" operator (i.e., the fitness consequences of having one or more genes of A mutated are the same), not just simply multiplying their fitness; similarly for "B". This is why no interaction genes also mean no modules allowed.

So how do you get what you want in this case? Enter the names of the modules in the epistasis component but have no term for ":" (the colon). Let's see an example:

```
fnme <- allFitnessEffects(epistasis = c("A" = 0.1,</pre>
                                           "B" = 0.2).
                           geneToModule = c("A" = "a1, a2",
                                              "B" = "b1, b2, b3")
evalAllGenotypes(fnme, order = FALSE, addwt = TRUE)
##
                 Genotype Fitness
## 1
                        WT
                               1.00
## 2
                        a1
                               1.10
## 3
                               1.10
                        a2
## 4
                        b1
                              1.20
## 5
                        b2
                              1.20
## 6
                              1.20
                        b3
## 7
                    a1, a2
                              1.10
```

```
## 8
                              1.32
                   a1, b1
## 9
                   a1, b2
                              1.32
## 10
                   a1, b3
                              1.32
## 11
                   a2, b1
                              1.32
## 12
                   a2, b2
                              1.32
## 13
                   a2, b3
                              1.32
                   b1, b2
                              1.20
## 14
                   b1, b3
                              1.20
## 15
## 16
                   b2, b3
                              1.20
## 17
               a1, a2, b1
                              1.32
## 18
               a1, a2, b2
                              1.32
               a1, a2, b3
                              1.32
## 19
## 20
               a1, b1, b2
                              1.32
               a1, b1, b3
## 21
                              1.32
## 22
               a1, b2, b3
                              1.32
## 23
               a2, b1, b2
                              1.32
               a2, b1, b3
## 24
                              1.32
## 25
               a2, b2, b3
                              1.32
## 26
               b1, b2, b3
                              1.20
## 27
          a1, a2, b1, b2
                              1.32
          a1, a2, b1, b3
## 28
                              1.32
## 29
          a1, a2, b2, b3
                              1.32
## 30
          a1, b1, b2, b3
                              1.32
## 31
          a2, b1, b2, b3
                              1.32
## 32 a1, a2, b1, b2, b3
                              1.32
```

In previous versions these was possible using the longer, still accepted way of specifying a : with a value of 0, but this is no longer needed:

```
fnme <- allFitnessEffects(epistasis = c("A" = 0.1,</pre>
                                           "B" = 0.2,
                                           "A : B'' = 0.0),
                            geneToModule = c("A" = "a1, a2",
                                              "B" = "b1, b2, b3"))
evalAllGenotypes(fnme, order = FALSE, addwt = TRUE)
##
                 Genotype Fitness
## 1
                        WT
                               1.00
## 2
                        a1
                               1.10
## 3
                        a2
                               1.10
## 4
                        b1
                               1.20
## 5
                        b2
                               1.20
## 6
                        b3
                               1.20
## 7
                    a1, a2
                               1.10
## 8
                    a1, b1
                               1.32
## 9
                    a1, b2
                               1.32
```

```
## 10
                               1.32
                   a1, b3
## 11
                   a2, b1
                               1.32
## 12
                   a2, b2
                              1.32
## 13
                   a2, b3
                              1.32
                   b1, b2
                              1.20
## 14
## 15
                   b1, b3
                              1.20
                   b2. b3
                              1.20
## 16
## 17
               a1, a2, b1
                              1.32
               a1, a2,
## 18
                       b2
                               1.32
## 19
               a1, a2, b3
                              1.32
## 20
               a1, b1, b2
                              1.32
## 21
               a1, b1, b3
                              1.32
## 22
               a1, b2, b3
                              1.32
## 23
               a2, b1, b2
                               1.32
## 24
               a2, b1, b3
                              1.32
## 25
               a2, b2, b3
                               1.32
## 26
               b1, b2, b3
                              1.20
## 27
           a1, a2, b1, b2
                              1.32
           a1, a2, b1, b3
## 28
                              1.32
## 29
           a1, a2, b2, b3
                               1.32
           a1, b1, b2, b3
## 30
                               1.32
## 31
           a2, b1, b2, b3
                               1.32
## 32 a1, a2, b1, b2, b3
                               1.32
```

This can, of course, be extended to more modules.

# 3.9 Synthetic viability

Synthetic viability and synthetic lethality (e.g., Ashworth, Lord, and Reis-Filho 2011; Hartman, Garvik, and Hartwell 2001) are just special cases of epistasis (section 3.7) but we deal with them here separately.

# 3.9.1 A simple synthetic viability example

A simple and extreme example of synthetic viability is shown in the following table, where the joint mutant has fitness larger than the wild type, but each single mutant is lethal.

Ā	В	Fitness
wt	wt	1
wt	M	0
Μ	wt	0
Μ	Μ	(1 + s)

where "wt" denotes wild type and "M" denotes mutant.

We can specify this (setting s = 0.2) as (I play around with spaces, to show there is a certain flexibility with them):

Now, let's look at all the genotypes (we use "addwt" to also get the wt, which by decree has fitness of 1), and disregard order:

Asking the program to consider the order of mutations of course makes no difference:

```
evalAllGenotypes(sv, order = TRUE, addwt = TRUE)
##
     Genotype Fitness
## 1
            WT
                    1.0
## 2
             \boldsymbol{A}
                    0.0
## 3
             B
                    0.0
## 4
        A > B
                    1.2
## 5 B > A
                    1.2
```

Another example of synthetic viability is shown in section 5.2.2.

Of course, if multiple simultaneous mutations are not possible in the simulations, it is not possible to go from the wildtype to the double mutant in this model where the single mutants are not viable.

#### 3.9.2 Synthetic viability, non-zero fitness, and modules

This is a slightly more elaborate case, where there is one module and the single mutants have different fitness between themselves, which is non-zero. Without the modules, this is the same as in Misra, Szczurek, and Vingron (2014), Figure 1b, which we go over in section 5.2.

A	В	Fitness
wt	wt	1
wt	${ m M}$	$1+s_b$
Μ	wt	$1+s_a$
M	M	$1 + s_{ab}$

where  $s_a, s_b < 0$  but  $s_{ab} > 0$ .

```
sa < -0.1
sb < -0.2
sab <- 0.25
sv2 <- allFitnessEffects(epistasis = c("-A : B" = sb,</pre>
                               "A : -B" = sa,
                               "A:B" = sab),
                          geneToModule = c(
                               "A" = "a1, a2",
                               "B" = "b"))
evalAllGenotypes(sv2, order = FALSE, addwt = TRUE)
##
      Genotype Fitness
## 1
             WT
                   1.00
## 2
                   0.90
             a1
## 3
             a2
                   0.90
## 4
              b
                   0.80
## 5
        a1, a2
                   0.90
## 6
         a1, b
                   1.25
## 7
         a2, b
                   1.25
## 8 a1, a2, b
                   1.25
```

And if we look at order, of course it makes no difference:

```
evalAllGenotypes(sv2, order = TRUE, addwt = TRUE)
##
         Genotype Fitness
## 1
                WT
                      1.00
## 2
                      0.90
                a1
                      0.90
## 3
                a2
## 4
                 b
                      0.80
## 5
          a1 > a2
                      0.90
## 6
           a1 > b
                      1.25
## 7
          a2 > a1
                      0.90
           a2 > b
## 8
                      1.25
## 9
           b > a1
                      1.25
## 10
           b > a2
                      1.25
## 11 a1 > a2 > b
                      1.25
## 12 a1 > b > a2
                      1.25
## 13 a2 > a1 > b
                      1.25
                      1.25
## 14 a2 > b > a1
## 15 b > a1 > a2
                      1.25
## 16 \ b > a2 > a1
                      1.25
```

# 3.10 Synthetic mortality or synthetic lethality

In contrast to section 3.9, here the joint mutant has decreased viability:

A	В	Fitness
A	В	Fitness
wt	wt	1
wt	M	$1+s_b$
M	wt	$1+s_a$
M	M	$1 + s_{ab}$

where  $s_a, s_b > 0$  but  $s_{ab} < 0$ .

```
sa <- 0.1
sb <- 0.2
sab < -0.8
sm1 <- allFitnessEffects(epistasis = c("-A : B" = sb,</pre>
                                "A : -B" = sa,
                                "A:B" = sab)
evalAllGenotypes(sm1, order = FALSE, addwt = TRUE)
##
     Genotype Fitness
## 1
            WT
                    1.0
## 2
             \boldsymbol{A}
                    1.1
             B
                    1.2
## 3
## 4
                    0.2
          A, B
```

And if we look at order, of course it makes no difference:

```
evalAllGenotypes(sm1, order = TRUE, addwt = TRUE)
     Genotype Fitness
##
## 1
            WT
                    1.0
## 2
             \boldsymbol{A}
                    1.1
## 3
             В
                    1.2
## 4
         A > B
                    0.2
                    0.2
## 5
        B > A
```

#### 3.11 Possible issues with Bozic model

#### 3.11.1 Synthetic viability using Bozic model

If we were to use the above specification with Bozic's models, we might not get what we think we should get:

What gives here? The simulation code would alert you of this (see section 3.11.2) in this particular case because there are "-1", which might indicate that this is not what you want. The problem is that you probably want the Death rate to be infinity (the birth rate was 0, so no clone viability, when we used birth rates —section 3.2.2).

Let us say so explicitly:

```
s < -0.2
svB <- allFitnessEffects(epistasis = c("-A : B" = -Inf,</pre>
                                          "A : -B" = -Inf,
                                          "A:B" = s)
evalAllGenotypes(svB, order = FALSE, addwt = TRUE, model = "Bozic")
     Genotype Death rate
## 1
            WT
                       1.0
## 2
             \boldsymbol{A}
                       Inf
## 3
             В
                       Inf
                       0.8
## 4
         A, B
```

Likewise, values of s larger than one have no effect beyond setting s = 1 (a single term of (1-1) will drive the product to 0, and as we cannot allow negative death rates negative values are set to 0):

```
s <- 1
svB1 <- allFitnessEffects(epistasis = c("-A : B" = -Inf,</pre>
                                           "A : -B" = -Inf,
                                           "A:B" = s)
evalAllGenotypes(svB1, order = FALSE, addwt = TRUE, model = "Bozic")
##
     Genotype Death rate
## 1
           WT
                         1
## 2
             \boldsymbol{A}
                       Inf
## 3
             B
                       Inf
## 4
        A, B
s <- 3
svB3 <- allFitnessEffects(epistasis = c("-A : B" = -Inf,</pre>
                                           "A : -B" = -Inf,
                                           "A:B" = s)
evalAllGenotypes(svB3, order = FALSE, addwt = TRUE, model = "Bozic")
     Genotype Death rate
##
            WT
## 1
## 2
             \boldsymbol{A}
                       Inf
             В
## 3
                       Inf
## 4
         A, B
                          0
```

Of course, death rates of 0.0 are likely to lead to trouble down the road, when we

actually conduct simulations (see section 3.11.2).

#### 3.11.2 Numerical issues with death rates of 0 in Bozic model

As we mentioned above (section 3.11.1) death rates of 0 can lead to trouble when using Bozic's model:

```
i1 <- allFitnessEffects(noIntGenes = c(1, 0.5))
evalAllGenotypes(i1, order = FALSE, addwt = TRUE,
                 model = "Bozic")
##
     Genotype Death_rate
## 1
           WT
## 2
                     0.0
            1
## 3
            2
                     0.5
                     0.0
## 4
         1, 2
i1_b <- oncoSimulIndiv(i1, model = "Bozic")</pre>
## Warning in nr_oncoSimul.internal(rFE = fp, birth = birth, death
## = death, : You are using a Bozic model with the new restriction
## specification, and you have at least one s of 1. If that gene is
## mutated, this will lead to a death rate of 0 and the simulations
## will abort when you get a non finite value.
##
##
   DEBUG2: Value of rnb = nan
##
   DEBUG2: Value of m = 1
##
##
   DEBUG2: Value of pe = 0
##
##
##
   DEBUG2: Value of pm = 0.9999999
##
##
   this is spP
##
## popSize = 1
## birth = 1
   death = 0
##
## W = 1
## R = 1
   mutation = 1e-06
##
## timeLastUpdate = 1096.35
   absfitness = -inf
##
   numMutablePos =1
##
##
   Unrecoverable exception: Algo 2: retval not finite. Aborting.
```

Of course, there is no problem in using the above with other models:

```
evalAllGenotypes(i1, order = FALSE, addwt = TRUE,
                 model = "Exp")
##
     Genotype Fitness
## 1
           WT
                   1.0
            1
## 2
                   2.0
            2
## 3
                  1.5
                  3.0
## 4
         1, 2
i1 e <- oncoSimulIndiv(i1, model = "Exp")</pre>
summary(i1 e)
##
     NumClones TotalPopSize LargestClone MaxNumDrivers MaxDriversLast
                                181588631
## 1
                   181659523
     NumDriversLargestPop TotalPresentDrivers FinalTime NumIter
##
## 1
                         0
                                              0
##
     HittedWallTime HittedMaxTries errorMF minDMratio minBMratio
## 1
              FALSE
                              FALSE
                                          NA
                                                  5e+05
                                                              5e+05
##
     OccurringDrivers
## 1
```

# 3.12 A longer example: Poset, epistasis, synthetic mortality and viability, order effects and genes without interactions, with some modules

We will now put together a complex example. We will use the poset from section 3.5.3 but will also add:

- Order effects that involve genes in the poset. In this case, if C happens before F, fitness decreases by 1-0.1. If it happens the other way around, there is no effect on fitness beyond their individual contributions.
- Order effects that involve two new modules, "H" and "I" (with genes "h1, h2" and "i1", respectively), so that if H happens before I fitness increases by 1+0.12.
- Synthetic mortality between modules "I" (already present in the epistatic interaction) and "J" (with genes "j1" and "j2"): the joint presence of these modules leads to cell death (fitness of 0).
- Synthetic viability between modules "K" and "M" (with genes "k1", "k2" and "m1", respectively), so that their joint presence is viable but adds nothing to fitness (i.e., mutation of both has fitness 1), whereas each single mutant has a fitness of 1-0.5.
- A set of 5 driver genes (n1, ..., n5) with fitness that comes from an exponential distribution with rate of 10.

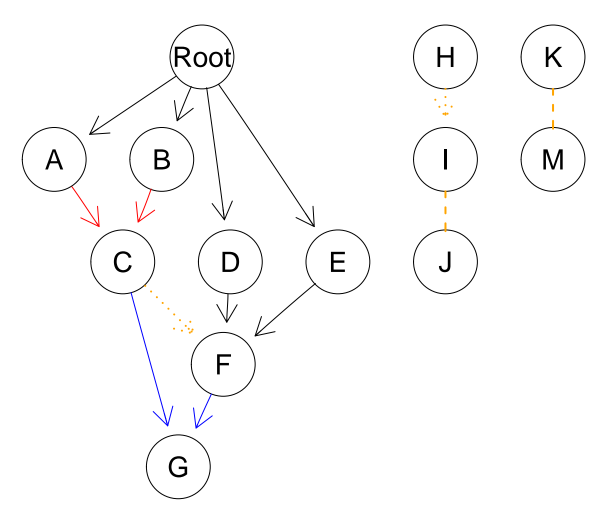
As we are specifying many different things, we will start by writing each set of effects separately:

```
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"),</pre>
```

```
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 \leftarrow c("-K:M" = -.5, "K:-M" = -.5)
epist <- c(sm, sv)</pre>
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 reproducibility
noint \leftarrow \text{rexp}(5, 10)
names(noint) <- paste0("n", 1:5)</pre>
fea <- allFitnessEffects(rT = p4, epistasis = epist,</pre>
                          orderEffects = oe,
                          noIntGenes = noint,
                          geneToModule = modules)
```

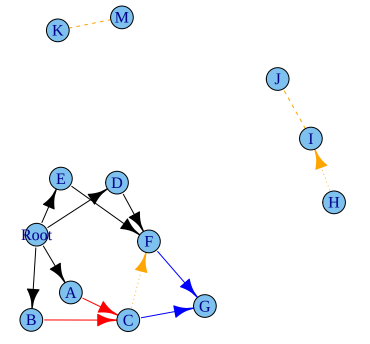
How does it look?

plot(fea)



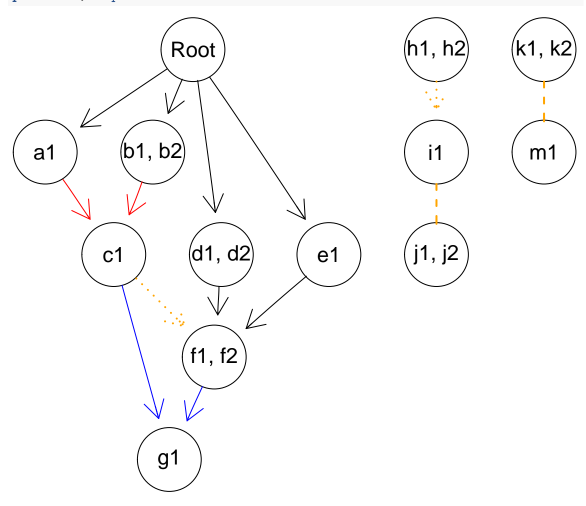
or

# plot(fea, "igraph")



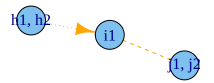
We can, if we want, expand the modules using a "graphNEL" graph

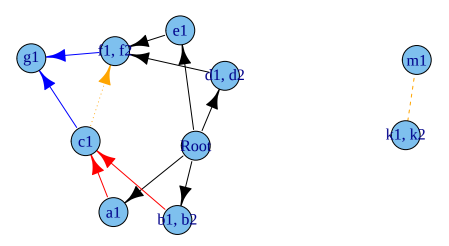
# plot(fea, expandModules = TRUE)



or an "igraph" one

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





We will not evaluate the fitness of all genotypes, since the number of all ordered genotypes is  $> 7 * 10^{22}$ . We will look at some specific genotypes:

```
evalGenotype("k1 > i1 > h2", fea) ## 0.5
## [1] 0.5
evalGenotype("k1 > h1 > i1", fea) ## 0.5 * 1.12
## [1] 0.56

evalGenotype("k2 > m1 > h1 > i1", fea) ## 1.12
## [1] 1.12

evalGenotype("k2 > m1 > h1 > i1 > c1 > n3 > f2", fea)
## [1] 0.005113
## 1.12 * 0.1 * (1 + noint[3]) * 0.05 * 0.9
```

Finally, let's generate some ordered genotypes randomly:

```
set.seed(2) ## for reproducibility
evalGenotype(randomGenotype(fea), fea, echo = TRUE, verbose = TRUE)
## Genotype: j2 > d2 > n4 > f1 > k2 > n1 > h2 > i1 > f2 > b1 > h1 > a1 > b2 > n3
   Individual s terms are : 0.0755182 0.118164 0.0145707 0.0139795 0.01 0.02 -0.
## Fitness: 0
## [1] 0
## Genotype: k2 > i1 > c1 > n1 > m1
## Individual s terms are : 0.0755182 -0.9
## Fitness: 0.107552
evalGenotype(randomGenotype(fea), fea, echo = TRUE, verbose = TRUE)
## Genotype: f2 > j1 > f1 > k1 > i1 > n4
## Individual s terms are : 0.0139795 -0.95 -1 -0.5
## Fitness: 0
## [1] 0
## Genotype: n2 > h1 > h2
## Individual s terms are : 0.118164
## Fitness: 1.11816
evalGenotype(randomGenotype(fea), fea, echo = TRUE, verbose = TRUE)
## Genotype: d2 > n1 > f2 > f1 > i1 > n5 > b1 > e1 > k2 > b2 > c1 > j1 > a1 > k1
## Individual s terms are : 0.0755182 0.0145707 0.0436069 0.01 0.02 -0.9 0.03 0.
## Fitness: 0
## [1] 0
## Genotype: d2 > k2 > c1 > f2 > n4 > m1 > n3 > f1 > b1 > g1 > n5 > h1 > j2
## Individual s terms are : 0.0145707 0.0139795 0.0436069 0.02 0.1 0.03 -0.95 0.
## Fitness: 0.0725829
evalGenotype(randomGenotype(fea), fea, echo = TRUE, verbose = TRUE)
## Genotype: a1 > m1 > f1 > c1 > i1 > d1 > b1 > n4 > d2 > n1 > e1 > k2 > j2 > n2
## Individual s terms are : 0.0755182 0.118164 0.0139795 0.01 0.02 -0.9 0.03 0.0.
## Fitness: 0
## [1] 0
## Genotype: h2 > c1 > f1 > n2 > b2 > a1 > n1 > i1
   Individual s terms are : 0.0755182 0.118164 0.01 0.02 -0.9 -0.95 -0.1 0.12
## Fitness: 0.00624418
evalGenotype(randomGenotype(fea), fea, echo = TRUE, verbose = TRUE)
## Genotype: g1 > j2 > m1 > d2 > n1 > n4 > i1 > b2 > f1
## Individual s terms are : 0.0755182 0.0139795 0.02 0.03 -0.95 0.3 -1 -0.5
## Fitness: 0
## [1] 0
## Genotype: h2 > j1 > m1 > d2 > i1 > b2 > k2 > d1 > b1 > n3 > n1 > q1 > h1 > c1
## Individual s terms are : 0.0755182 0.0145707 0.0436069 0.01 0.02 -0.9 0.03 0.
evalGenotype(randomGenotype(fea), fea, echo = TRUE, verbose = TRUE)
## Genotype: i1 > k2 > d1 > d2 > n4 > f2 > j2 > c1 > a1 > j1 > n1 > n3 > h1 > m1
## Individual s terms are : 0.0755182 0.118164 0.0145707 0.0139795 0.0436069 0.0
```

```
## [1] 0
## Genotype: n1 > m1 > n3 > i1 > j1 > n5 > k1
## Individual s terms are : 0.0755182 0.0145707 0.0436069 -1
## Fitness: 0
evalGenotype(randomGenotype(fea), fea, echo = TRUE, verbose = TRUE)
## Genotype: n2 > a1 > c1 > d2 > j1 > e1 > k1 > b2 > d1 > n3 > j2 > f2 > i1 > g1
   Individual s terms are : 0.0755182 0.118164 0.0145707 0.0139795 0.0436069 0.0
## Fitness: 0
## [1] 0
## Genotype: d2 > n1 > g1 > f1 > f2 > c1 > b1 > d1 > k1 > a1 > b2 > i1 > n4 > h2
## Individual s terms are : 0.0755182 0.118164 0.0139795 0.01 0.02 -0.9 0.03 -0.
  Fitness: 0.00420528
evalGenotype(randomGenotype(fea), fea, echo = TRUE, verbose = TRUE)
## Genotype: d2 > a1 > h2
  Individual s terms are : 0.01 0.03
## Fitness: 1.04
## [1] 1.04
## Genotype: j1 > f1 > j2 > a1 > n4 > c1 > n3 > k1 > d1 > h1
   Individual s terms are : 0.0145707 0.0139795 0.01 0.1 0.03 -0.95 -0.5
## Fitness: 0.0294308
evalGenotype(randomGenotype(fea), fea, echo = TRUE, verbose = TRUE)
## Genotype: n2 > f2
## Individual s terms are : 0.118164 -0.95
## Fitness: 0.05591
## [1] 0.05591
## Genotype: n5 > f2 > f1 > h2 > n4 > c1 > n3 > b1
   Individual s terms are: 0.0145707 0.0139795 0.0436069 0.02 0.1 -0.95
## Fitness: 0.0602298
evalGenotype(randomGenotype(fea), fea, echo = TRUE, verbose = TRUE)
## Genotype: n5 > n1 > d1 > f1 > c1 > b1 > n2 > i1 > a1 > n3 > n4 > e1 > k2 > b2
## Individual s terms are : 0.0755182 0.118164 0.0145707 0.0139795 0.0436069 0.0
## Fitness: 0
## [1] 0
## Genotype: h1 > d1 > f2
## Individual s terms are : 0.03 -0.95
## Fitness: 0.0515
```

# 3.13 Homozygosity, heterozygosity, oncogenes, tumor suppressors

We are using what is conceptually a single linear chromosome. However, you can use it to model scenarios where the numbers of copies affected matter, by properly duplicating the genes.

Suppose we have a tumor suppressor gene, G, with two copies, one from Mom and one from Dad. We can have a table like:

$\overline{O_M}$	$O_D$	Fitness
wt	wt	1
wt	${ m M}$	1
M	wt	1
M	M	(1+s)

where s > 0, meaning that you need two hits, one in each copy, to trigger the clonal expansion.

What about oncogenes? A simple model is that one single hit leads to clonal expansion and additional hits lead to no additional changes, as in this table for gene O, where again the M or D subscript denotes the copy from Mom or from Dad:

$\overline{O_M}$	$O_D$	Fitness
wt	wt	1
wt	${ m M}$	(1+s)
M	wt	(1+s)
M	M	(1+s)

If you have multiple copies you can proceed similarly. As you can see, these are nothing but special cases of synthetic mortality (3.10), synthetic viability (3.9) and epistasis (3.7).

# 3.14 Gene-specific mutation rates

You can specify gene-specific mutation rates. Instead of passing a scalar value for mu, you pass a named vector. (This does not work with the old v. 1 format, though; yet another reason to stop using that format). This is a simple example (many more are available in the tests, see file ./tests/testthat/test.per-gene-mutation-rates.R).

# 3.15 Mutator genes

You can specify mutator/antimutator genes (e.g. Gerrish et al. 2007; Tomlinson, Novelli, and Bodmer 1996). These are genes that, when mutated, lead to an

increase/decrease in the mutation rate all over the genome (similar to what happens with, say, mutations in mismatch-repair genes or microsatellite instability in cancer).

The specification is very similar to that for fitness effects, except we do not (at least for now) allow the use of DAGs nor of order effects (we have seen no reference in the literature to suggest any of these would be relevant). You can, however, specify epistasis and use modules. Note that the mutator genes must be a subset of the genes in the fitness effects; if you want to have mutator genes that have no direct fitness effects, give them a fitness effect of 0.

This first is a very simple example with simple fitness effects and modules for mutators. We will specify the fitness and mutator effects and evaluate the fitness and mutator effects:

```
fe2 <- allFitnessEffects(noIntGenes =</pre>
                         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,</pre>
                                        "B" = 10.
                                        "C" = 3),
                         geneToModule = c("A" = "a1, a2",
                                           "B" = "b1, b2, b3",
                                           "C" = "c1, c2"))
## Show the fitness effect of a specific genotype
evalGenotype("a1, c2", fe2, verbose = TRUE)
##
## Individual s terms are : 0.1 -0.2
## [1] 0.88
## Show the mutator effect of a specific genotype
evalGenotypeMut("a1, c2", fm2, verbose = TRUE)
##
## Individual mutator product terms are : 5 3
## [1] 15
## Fitness and mutator of a specific genotype
evalGenotypeFitAndMut("a1, c2", fe2, fm2, verbose = TRUE)
##
##
    Individual s terms are : 0.1 -0.2
##
    Individual mutator product terms are : 5 3
## [1] 0.88 15.00
```

You can also use the evalAll functions. We do not show the output here to avoid cluttering the vignette:

```
## 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)
```

Building upon the above, the next is an example where we have a bunch of no interaction genes that affect fitness, and a small set of genes that affect the mutation rate (but have no fitness effects).

Let us check what the effects are of a few genotypes:

```
## These only affect mutation, not fitness
evalGenotypeFitAndMut("a1, a2", fe3, fm3, verbose = TRUE)
##
   Individual s terms are : 0 0
##
##
## Individual mutator product terms are : 5
## [1] 1 5
evalGenotypeFitAndMut("a1, b3", fe3, fm3, verbose = TRUE)
##
   Individual s terms are : 0 0
##
##
## Individual mutator product terms are : 5 10
## [1] 1 50
## These only affect fitness: the mutator multiplier is 1
evalGenotypeFitAndMut("g1", fe3, fm3, verbose = TRUE)
```

```
##
##
   Individual s terms are : 0.019206
## [1] 1.019 1.000
evalGenotypeFitAndMut("g3, g9", fe3, fm3, verbose = TRUE)
##
   Individual s terms are : 0.0530139 0.0592025
##
## [1] 1.115 1.000
## These affect both
evalGenotypeFitAndMut("g3, g9, a2, b3", fe3, fm3, verbose = TRUE)
##
##
   Individual s terms are : 0 0 0.0530139 0.0592025
##
##
   Individual mutator product terms are : 5 10
## [1] 1.115 50.000
```

Finally, we will do a simulation with those data

```
## We do not show this in the vignette to avoid cluttering it
## with output
mue1
```

Of course, it is up to you to keep things reasonable: mutator effects are multiplicative, so if you specify, say, 20 genes (without modules), or 20 modules, each with a mutator effect of 50, the overall mutation rate can be increased by a factor of  $50^{20}$  and that is unlikely to be what you really want (see also section 14.7).

You can play with the following case (an extension of the example above), where a clone with a mutator phenotype and some fitness enhancing mutations starts giving rise to many other clones, some with additional mutator effects, and thus leading to the number of clones blowing up (as some also accumulate additional fitness-enhancing mutations). Things start getting out of hand shortly after time 250. The code below takes a few minutes to run and is not executed here, but you can run it to get an idea of the increase in the number of clones and their relationships (the usage of plotClonePhylog is explained in section 8).

```
set.seed(1) ## for reproducibility
## 17 genes, 7 with no direct fitness effects
ni <- c(rep(0, 7), runif(10, min = -0.01, max = 0.1))</pre>
```

```
names(ni) <- c("a1", "a2", "b1", "b2", "b3", "c1", "c2",</pre>
               paste0("g", 1:10))
## Next is for nicer figure labeling.
## Consider as drivers genes with s >0
gp <- which(ni > 0)
fe3 <- allFitnessEffects(noIntGenes = ni,</pre>
                          drvNames = names(ni)[gp])
set.seed(12)
mue1 <- oncoSimulIndiv(fe3, muEF = fm3,</pre>
                       mu = 1e-6,
                       initSize = 1e5,
                       model = "McFL",
                       detectionSize = 5e6,
                       finalTime = 270,
                       keepPhylog = TRUE,
                        onlyCancer = FALSE)
mue1
## If you decrease N even further it gets even more cluttered
op <- par(ask = TRUE)
plotClonePhylog(mue1, N = 10, timeEvents = TRUE)
plot(mue1, plotDrivers = TRUE, addtot = TRUE,
     plotDiversity = TRUE)
## The stacked plot is slow; be patient
## Most clones have tiny population sizes, and their lines
## are piled on top of each other
plot(mue1, addtot = TRUE,
     plotDiversity = TRUE, type = "stacked")
par(op)
```

# 4 Plotting fitness landscapes

The evalAllGenotypes and related functions allow you to obtain tables of the genotype to fitness mappings. It might be more convenient to actually plot that, allowing us to quickly identify local minima and maxima and get an idea of how the fitness landscape looks.

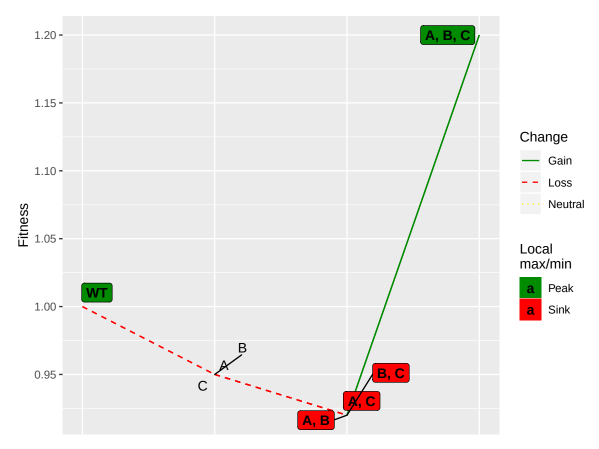
In plotFitnessLandscape I have blatantly and shamelessly copied most of the looks of the plots of MAGELLAN (Brouillet et al. 2015) (see also http://wwwabi.snv.jussieu.fr/public/Magellan/), a very nice web-based tool for fitness landscape plotting and analysis (MAGELLAN provides some other extra functionality and epistasis statistics not provided here).

As an example, let us show the example of Weissman et al. we saw in 5.4:

```
d1 <- -0.05 ## single mutant fitness 0.95
d2 <- -0.08 ## double mutant fitness 0.92
d3 <- 0.2 ## triple mutant fitness 1.2
s2 <- ((1 + d2)/(1 + d1)^2) - 1
s3 <- ( (1 + d3)/((1 + d1)^3 * (1 + s2)^3) ) - 1

wb <- allFitnessEffects(
    epistasis = c(
        "A" = d1,
        "B" = d1,
        "C" = d1,
        "A:B" = s2,
        "A:C" = s2,
        "B:C" = s2,
        "A:B:C" = s3))</pre>
```

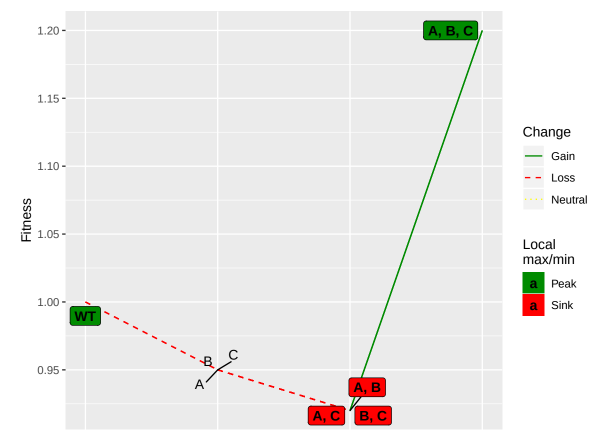
plotFitnessLandscape(wb, use ggrepel = TRUE)



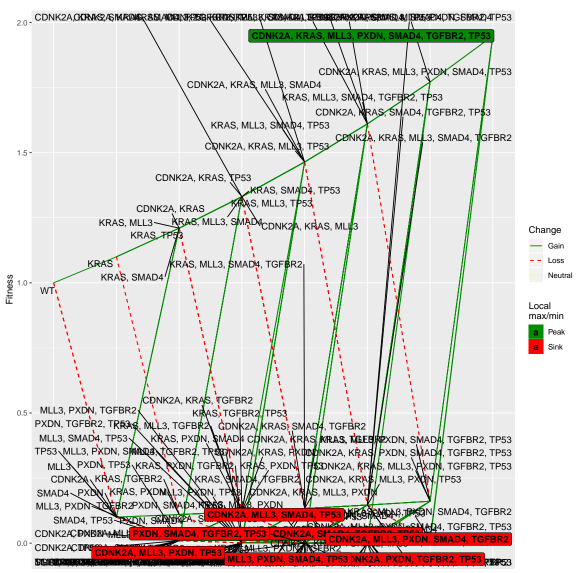
We have set  ${\tt use\_ggrepel}$  = TRUE to avoid overlap of labels.

For some types of objects, directly invoking plot will give you the fitness landscape plot:

```
(ewb <- evalAllGenotypes(wb, order = FALSE))</pre>
     Genotype Fitness
            \boldsymbol{A}
                  0.95
## 1
## 2
            B
                  0.95
## 3
          C
                  0.95
       A, B
                 0.92
## 4
## 5
         A, C
                  0.92
## 6
       B, C
                  0.92
## 7 A, B, C
                  1.20
plot(ewb, use_ggrepel = TRUE)
```



This is example (section 5.5) will give a very busy plot:



# 5 Specifying fitness effects: some examples from the literature

# 5.1 Bauer et al., 2014

In the model of Bauer and collaborators (Bauer, Siebert, and Traulsen 2014, 54) we have "For cells without the primary driver mutation, each secondary driver mutation leads to a change in the cell's fitness by  $s_P$ . For cells with the primary driver mutation, the fitness advantage obtained with each secondary driver mutation is  $s_{DP}$ ."

The proliferation probability is given as:

- $\frac{1}{2}(1+s_p)^k$  when there are k secondary drivers mutated and no primary diver;
- $\frac{1}{2} \frac{1+S_D^+}{1+S_D^-} (1+S_{DP})^k$  when the primary driver is mutated;

apoptosis is one minus the proliferation rate.

# 5.1.1 Using a DAG

We cannot find a simple mapping from their expressions to our fitness parameterization, but we can get fairly close by using a DAG; in this one, note the unusual feature of having one of the "s" terms (that for the driver dependency on root) be negative. Using the parameters given in the legend of their Figure 3 for  $s_p, S_D^+, S_D^-, S_{DP}$  and obtaining that negative value for the dependency of the driver on root we can do:

```
K <- 4
sp < -1e-5
sdp <- 0.015
sdplus <- 0.05
sdminus <- 0.1
cnt \langle -(1 + sdplus)/(1 + sdminus)
prod_cnt <- cnt - 1</pre>
bauer <- data.frame(parent = c("Root", rep("D", K)),</pre>
                     child = c("D", paste0("s", 1:K)),
                     s = c(prod_cnt, rep(sdp, K)),
                     sh = c(0, rep(sp, K)),
                     typeDep = "MN")
fbauer <- allFitnessEffects(bauer)</pre>
(b1 <- evalAllGenotypes(fbauer, order = FALSE, addwt = TRUE))
##
                Genotype Fitness
                       WT 1.0000
## 1
## 2
                           0.9545
## 3
                       s1
                           1.0000
                       s2
                          1.0000
## 4
## 5
                       s3 1.0000
## 6
                       s4 1.0000
## 7
                   D, s1 0.9689
```

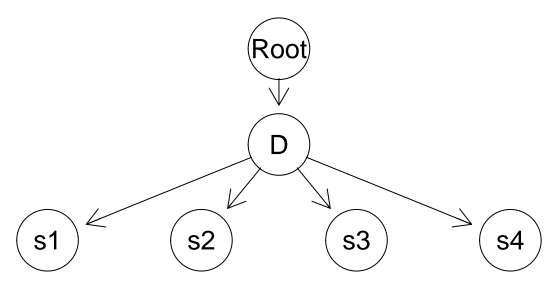
```
## 8
                   D, s2
                           0.9689
## 9
                   D, s3
                           0.9689
## 10
                   D, s4
                           0.9689
## 11
                  s1, s2
                           1.0000
                  s1, s3
                           1.0000
## 12
## 13
                  s1, s4
                           1.0000
## 14
                  s2, s3
                           1.0000
## 15
                  s2, s4
                           1.0000
## 16
                  s3, s4
                           1.0000
              D, s1, s2
## 17
                           0.9834
## 18
              D, s1, s3
                           0.9834
              D, s1, s4
## 19
                           0.9834
## 20
              D, s2, s3
                           0.9834
              D, s2, s4
## 21
                           0.9834
## 22
              D, s3, s4
                           0.9834
             s1, s2, s3
## 23
                           1.0000
## 24
             s1, s2, s4
                           1.0000
## 25
             s1, s3, s4
                           1.0000
             s2, s3, s4
## 26
                           1.0000
          D, s1, s2, s3
## 27
                           0.9981
          D, s1, s2, s4
## 28
                           0.9981
          D, s1, s3, s4
## 29
                           0.9981
## 30
          D, s2, s3, s4
                           0.9981
## 31
         s1, s2, s3, s4
                           1.0000
## 32 D, s1, s2, s3, s4
                           1.0131
```

(We use "D" for "driver" or "primary driver", as is it is called in the original paper, and "s" for secondary drivers, somewhat similar to passengers).

Note that what we specify as "typeDep" is irrelevant (MN, SMN, or XMPN make no difference).

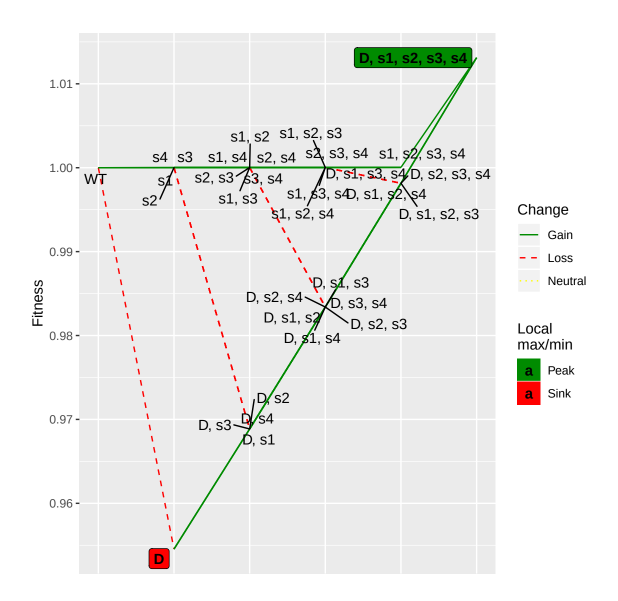
This is the DAG:

```
plot(fbauer)
```



And if you compare the tabular output of evalAllGenotypes you can see that the values of fitness reproduces the fitness landscape that they show in their Figure 1. We can also use our plot for fitness landscapes:

plot(b1, use\_ggrepel = TRUE)



#### 5.1.2 Specifying fitness of genotypes directly

An alternative approach to specify the fitness, if the number of genotypes is reasonably small, is to directly evaluate fitness as given by their expressions. Then, use the genotFitness argument to allFitnessEffects.

We will create all possible genotypes; then we will write a function that gives the fitness of each genotype according to their expression; finally, we will call this function on the data frame of genotypes, and pass this data frame to allFitnessEffects.

Now, show the fitness of all genotypes:

```
evalAllGenotypes(bauer2, order = FALSE, addwt = TRUE)
##
              Genotype Fitness
## 1
                    WT 1.0000
## 2
                     D 0.9545
## 3
                    s1 1.0000
## 4
                    s2 1.0000
## 5
                    s3 1.0000
## 6
                    s4 1.0000
                 D, s1 0.9689
## 7
## 8
                 D, s2 0.9689
## 9
                 D, s3 0.9689
## 10
                 D, s4 0.9689
## 11
                s1, s2 1.0000
## 12
                s1, s3 1.0000
## 13
                s1, s4 1.0000
                s2, s3 1.0000
## 14
## 15
                s2, s4 1.0000
## 16
                s3, s4 1.0000
## 17
             D, s1, s2 0.9834
## 18
             D, s1, s3 0.9834
## 19
             D, s1, s4 0.9834
## 20
             D, s2, s3 0.9834
## 21
             D, s2, s4 0.9834
## 22
             D, s3, s4 0.9834
            s1, s2, s3 1.0000
## 23
## 24
            s1, s2, s4 1.0000
## 25
            s1, s3, s4
                       1.0000
## 26
            s2, s3, s4 1.0000
```

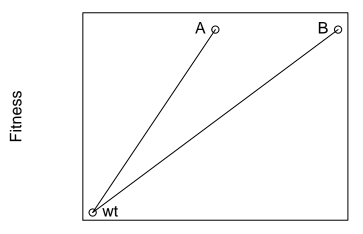
```
## 27
       D, s1, s2, s3
                          0.9981
## 28
          D, s1, s2, s4
                          0.9981
## 29
          D, s1, s3, s4
                          0.9981
          D, s2, s3, s4
## 30
                          0.9981
## 31
         s1, s2, s3, s4
                          1.0000
## 32 D, s1, s2, s3, s4
                          1.0131
```

Can we use modules in this example, if we use the "lego system"? Sure, as in any other case.

# 5.2 Misra et al., 2014

Figure 1 of Misra, Szczurek, and Vingron (2014) presents three scenarios which are different types of epistasis.

# 5.2.1 Example 1.a



In that figure it is evident that the fitness effect of "A" and "B" are the same. There are two different models depending on whether "AB" is just the product of both, or there is epistasis. In the first case probably the simplest is:

```
s <- 0.1 ## or whatever number
m1a1 <- allFitnessEffects(data.frame(parent = c("Root", "Root"),</pre>
                                         child = c("A", "B"),
                                         s = s,
                                         sh = 0,
                                         typeDep = "MN"))
evalAllGenotypes(m1a1, order = FALSE, addwt = TRUE)
##
      Genotype Fitness
## 1
            WT
                   1.00
## 2
             \boldsymbol{A}
                   1.10
## 3
             В
                   1.10
          A, B
                   1.21
```

If the double mutant shows epistasis, as we saw before (section 3.7.1) we have a range of options. For example:

```
s < -0.1
sab <- 0.3
m1a2 <- allFitnessEffects(epistasis = c("A:-B" = s,</pre>
                                             "-A:B" = s,
                                             "A:B" = sab))
evalAllGenotypes(m1a2, order = FALSE, addwt = TRUE)
      Genotype Fitness
##
            WT
## 1
                    1.0
                    1.1
## 2
             \boldsymbol{A}
## 3
             B
                    1.1
## 4
      A, B
                    1.3
```

But we could also modify the graph dependency structure, and we have to change the value of the coefficient, since that is what multiplies each of the terms for "A" and "B":  $(1 + s_{AB}) = (1 + s)^2(1 + s_{AB3})$ 

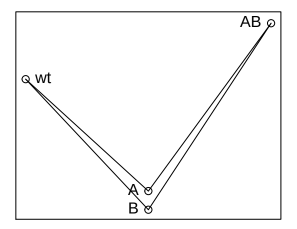
```
sab3 \leftarrow ((1 + sab)/((1 + s)^2)) - 1
m1a3 <- allFitnessEffects(data.frame(parent = c("Root", "Root"),</pre>
                                         child = c("A", "B"),
                                         s = s,
                                         sh = 0,
                                         typeDep = "MN"),
                            epistasis = c("A:B" = sab3))
evalAllGenotypes(m1a3, order = FALSE, addwt = TRUE)
##
     Genotype Fitness
## 1
            WT
                    1.0
## 2
                    1.1
             \boldsymbol{A}
## 3
             B
                    1.1
## 4
       A, B
                    1.3
```

And, obviously

#### **5.2.2** Example 1.b

This is a specific case of synthetic viability (see also section 3.9):

itness



Here,  $S_A, S_B < 0$ ,  $S_B < 0$ ,  $S_{AB} > 0$  and  $(1 + S_{AB})(1 + S_A)(1 + S_B) > 1$ .

As before, we can specify this in several different ways. The simplest is to specify all genotypes:

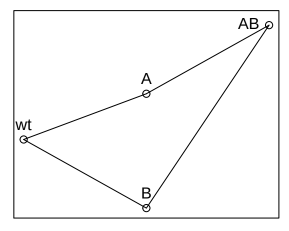
```
sa < -0.6
sb < -0.7
sab <- 0.3
m1b1 <- allFitnessEffects(epistasis = c("A:-B" = sa,
                                          "-A:B" = sb,
                                          "A:B" = sab))
evalAllGenotypes(m1b1, order = FALSE, addwt = TRUE)
##
     Genotype Fitness
## 1
            WT
                   1.0
## 2
             Α
                   0.4
                   0.3
             В
## 3
         A, B
                   1.3
## 4
```

We could also use a tree and modify the "sab" for the epistasis, as before (5.2.1).

# **5.2.3** Example 1.c

The final case, in figure 1.c of Misra et al., is just epistasis, where a mutation in one of the genes is deleterious (possibly only mildly), in the other is beneficial, and the double mutation has fitness larger than any of the other two.

ness



```
Here we have that s_A > 0, s_B < 0, (1 + s_{AB})(1 + s_A)(1 + s_B) > (1 + s_{AB}) so s_{AB} > \frac{-s_B}{1+s_B}
```

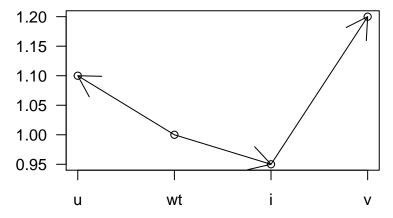
As before, we can specify this in several different ways. The simplest is to specify all genotypes:

```
sa <- 0.2
sb < -0.3
sab <- 0.5
m1c1 <- allFitnessEffects(epistasis = c("A:-B" = sa,</pre>
                                             "-A:B" = sb,
                                             "A:B" = sab))
evalAllGenotypes(m1c1, order = FALSE, addwt = TRUE)
##
      Genotype Fitness
## 1
            WT
                     1.0
## 2
              \boldsymbol{A}
                     1.2
## 3
              В
                     0.7
                     1.5
## 4
          A, B
```

We could also use a tree and modify the "sab" for the epistasis, as before (5.2.1).

# 5.3 Ochs and Desai, 2015

In Ochs and Desai (2015) the authors present a model shown graphically as (the actual numerical values are arbitrarily set by me):



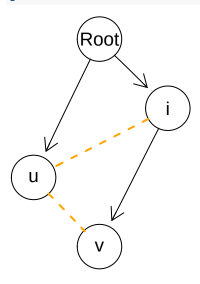
In their model,  $s_u > 0$ ,  $s_v > s_u$ ,  $s_i < 0$ , we can only arrive at v from i, and the mutants "ui" and "uv" can never appear as their fitness is 0, or  $-\infty$ , so  $s_{ui} = s_{uv} = -1$  (or  $-\infty$ ).

We can specify this combining a graph and epistasis specifications:

```
su <- 0.1
si <- -0.05
fvi <- 1.2 ## the fitness of the vi mutant
sv <- (fvi/(1 + si)) - 1
sui <- suv <- -1
od <- allFitnessEffects(</pre>
```

A figure showing that model is

#### plot(od)



And the fitness of all genotype is

```
evalAllGenotypes(od, order = FALSE, addwt = TRUE)
     Genotype Fitness
## 1
          WT
                 1.00
## 2
           i
                 0.95
## 3
                1.10
           u
## 4
                 0.00
           v
## 5
        i, u
                0.00
## 6
        i, v
                 1.20
## 7
                 0.00
         u, v
## 8 i, u, v
                 0.00
```

We could alternatively have specified fitness either directly specifying the fitness of each genotype or specifying epistatic effects. Let us use the second approach:

%% this was wrong %% u <- 0.2; i <- -0.02; vi <- 0.6; ui <- uv <- -Inf

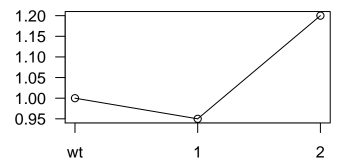
```
"v:-i" = -Inf, "v:i" = vi))
evalAllGenotypes(od2, addwt = TRUE)
     Genotype Fitness
## 1
            WT
                   1.00
                   0.95
## 2
             i
## 3
                   1.10
             u
## 4
                   0.00
             υ
## 5
                   0.00
          i, u
## 6
                   1.20
          i, v
                   0.00
                   0.00
      i, u, v
```

We will return to this model when we explain the usage of fixation for stopping the simulations (see 6.3.3 and 6.3.4).

# 5.4 Weissman et al., 2009

In their figure 1a, Weissman et al. (2009) present this model (actual numeric values are set arbitrarily)

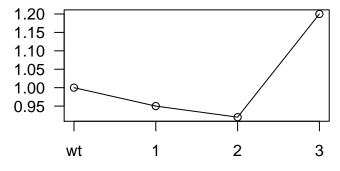
### 5.4.1 Figure 1.a



where the "1" and "2" in the figure refer to the total number of mutations in two different loci. This is, therefore, very similar to the example in section 5.2.2. Here we have, in their notation,  $\delta_1 < 0$ , fitness of single "A" or single "B" =  $1 + \delta_1$ ,  $S_{AB} > 0$ ,  $(1 + S_{AB})(1 + \delta_1)^2 > 1$ .

#### **5.4.2** Figure 1.b

In their figure 1b they show



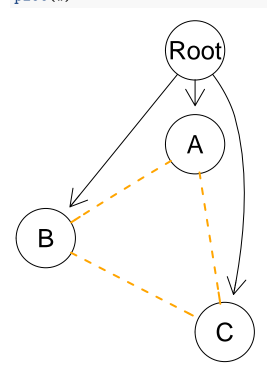
Where, as before, 1, 2, 3, denote the total number of mutations over three different loci and  $\delta_1 < 0$ ,  $\delta_2 < 0$ , fitness of single mutant is  $(1 + \delta_1)$ , of double mutant is  $(1 + \delta_2)$  so that  $(1 + \delta_2) = (1 + \delta_1)^2(1 + s_2)$  and of triple mutant is  $(1 + \delta_3)$ , so that  $(1 + \delta_3) = (1 + \delta_1)^3(1 + s_2)^3(1 + s_3)$ .

We can specify this combining a graph with epistasis:

```
d1 <- -0.05 ## single mutant fitness 0.95
d2 <- -0.08 ## double mutant fitness 0.92
d3 <- 0.2
           ## triple mutant fitness 1.2
s2 \leftarrow ((1 + d2)/(1 + d1)^2) - 1
s3 \leftarrow ((1 + d3)/((1 + d1)^3 * (1 + s2)^3)) - 1
w <- allFitnessEffects(</pre>
    data.frame(parent = c("Root", "Root", "Root"),
                child = c("A", "B", "C"),
                s = d1,
                sh = -1,
                typeDep = "MN"),
    epistasis = c(
        "A:B" = s2,
        "A:C" = s2,
        "B:C" = s2,
        "A:B:C" = s3))
```

The model can be shown graphically as:

#### plot(w)



And fitness of all genotypes is:

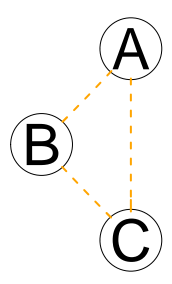
```
evalAllGenotypes(w, order = FALSE, addwt = TRUE)
     Genotype Fitness
## 1
            WT
                   1.00
## 2
             \boldsymbol{A}
                   0.95
## 3
             B
                   0.95
             C
## 4
                   0.95
## 5
          A, B
                   0.92
## 6
          A, C
                   0.92
## 7
          B, C
                   0.92
## 8 A, B, C
                   1.20
```

Alternatively, we can directly specify what each genotype adds to the fitness, given the included genotype. This is basically replacing the graph by giving each of "A", "B", and "C" directly:

```
wb <- allFitnessEffects(</pre>
    epistasis = c(
         ^{"}A" = d1,
         "B" = d1,
         "C" = d1,
         "A:B" = s2,
         "A:C" = s2,
         "B:C" = s2,
         "A:B:C" = s3))
evalAllGenotypes(wb, order = FALSE, addwt = TRUE)
##
      Genotype Fitness
## 1
            WT
                   1.00
## 2
             \boldsymbol{A}
                    0.95
## 3
             В
                    0.95
              C
                   0.95
## 4
          A, B
## 5
                   0.92
## 6
          A, C
                    0.92
## 7
          B, C
                    0.92
## 8 A, B, C
                   1.20
```

The plot, of course, is not very revealing and we cannot show that there is a three-way interaction (only all three two-way interactions):

```
plot(wb)
```



As we have seen several times already (sections 3.7.1, 3.7.2, 3.7.3) we can also give the genotypes directly and, consequently, the fitness of each genotype (not the added contribution):

```
wc <- allFitnessEffects(</pre>
    epistasis = c(
         "A:-B:-C" = d1,
         "B:-C:-A" = d1,
         "C:-A:-B" = d1,
         "A:B:-C" = d2,
         "A:C:-B" = d2,
         "B:C:-A" = d2,
         "A:B:C" = d3))
evalAllGenotypes(wc, order = FALSE, addwt = TRUE)
##
     Genotype Fitness
## 1
           WT
                   1.00
## 2
             \boldsymbol{A}
                   0.95
## 3
             B
                   0.95
             C
                   0.95
## 4
          A, B
## 5
                   0.92
## 6
          A. C
                   0.92
## 7
          B, C
                   0.92
## 8 A, B, C
                   1.20
```

## 5.5 Gerstung et al., 2011, pancreatic cancer poset

Similar to what we did in v.1 (see section 15.1) we can specify the pancreatic cancer poset in Gerstung et al. (2011) (their figure 2B, left). We use directly the names of the genes, since that is immediately supported by the new version.

```
"TP53", "TP53", "MLL3"),

child = c("KRAS", "SMAD4", "CDNK2A",

"TP53", "MLL3",

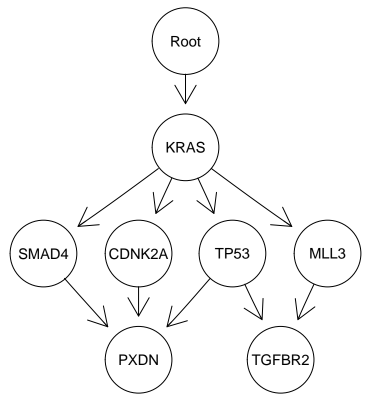
rep("PXDN", 3), rep("TGFBR2", 2)),

s = 0.1,

sh = -0.9,

typeDep = "MN"))

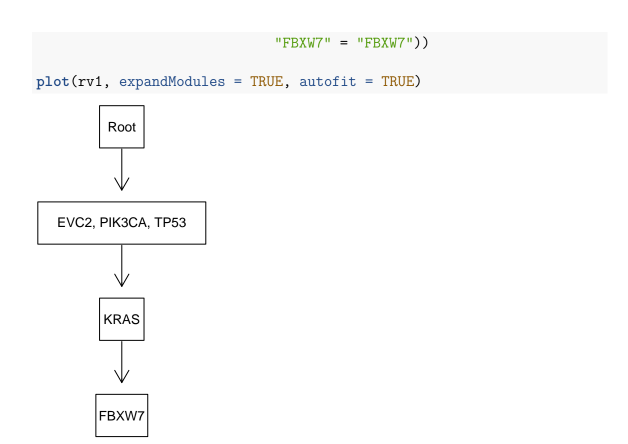
plot(pancr)
```



Of course the "s" and "sh" are set arbitrarily here.

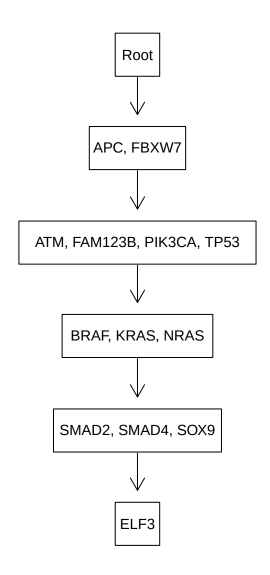
## 5.6 Raphael and Vandin's 2014 modules

In Raphael and Vandin (2015), the authors show several progression models in terms of modules. We can code the extended poset for the colorectal cancer model in their Figure 4.a is (s and sh are arbitrary):



We have used the (experimental) autofit option to fit the labels to the edges. Note how we can use the same name for genes and modules, but we need to specify all the modules.

Their Figure 5b is



# 6 Running and plotting the simulations: starting, ending, and examples

### 6.1 Starting and ending

After you have decided the specifics of the fitness effects and the model, you need to decide:

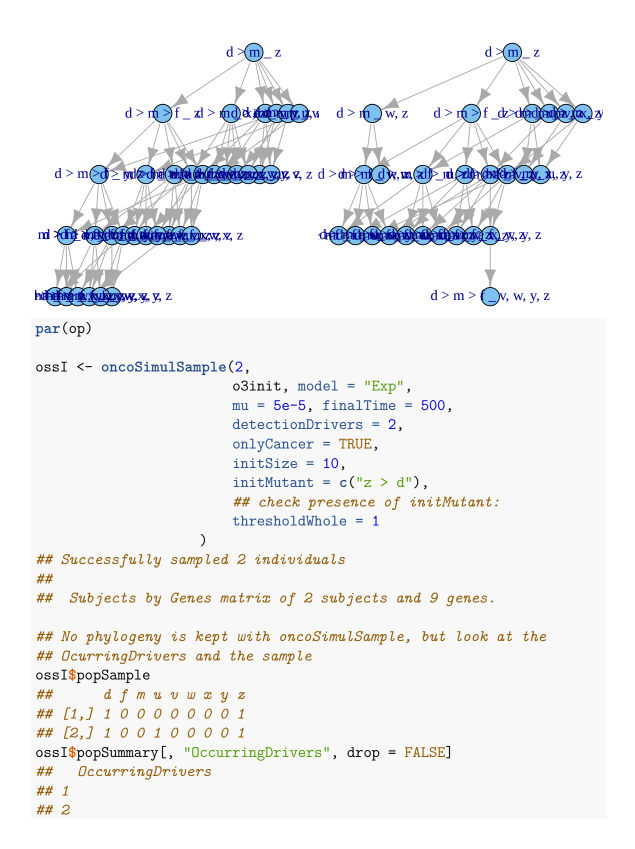
- Where will you start your simulation from. This involves deciding the initial population size (argument initSize) and, possibly, the genotype of the initial population; the later is covered in section 6.2.
- When will you stop it: how long to run it, and whether or not to require simulations to reach cancer (under some definition of what it means to reach cancer). This is covered in 6.3.

### 6.2 Can I start the simulation from a specific mutant?

You bet. In version 2 you can specify the genotype for the initial mutant with the same flexibility as in evalGenotype. Here we show a couple of examples (we use the representation of the parent-child relationships —discussed in section 8— of the clones so that you can see which clones appear, and from which, and check that we are not making mistakes).

```
o3init <- allFitnessEffects(orderEffects = c(
                                 "M > D > F" = 0.99,
                                 "D > M > F" = 0.2,
                                 "D > M"
                                               = 0.1,
                                 "M > D"
                                               = 0.9),
                            noIntGenes = c("u" = 0.01,
                                              "v" = 0.01.
                                              w'' = 0.001
                                              "x" = 0.0001,
                                              "v" = -0.0001.
                                              z'' = -0.001,
                            geneToModule =
                                 c("M" = "m",
                                   ^{\prime\prime}F^{\prime\prime} = ^{\prime\prime}f^{\prime\prime}.
                                   "D" = "d")
oneI <- oncoSimulIndiv(o3init, model = "McFL",</pre>
                           mu = 5e-5, finalTime = 500,
                           detectionDrivers = 3,
                           onlyCancer = FALSE,
                           initSize = 1000,
                           keepPhylog = TRUE,
                           initMutant = c("m > u > d")
```

```
plotClonePhylog(oneI, N = 0)
                                              m > d
                         [u, z m > d(> f)]
                                       _m, wd(mf
m > h \gg f(\underline{1} > u)(\underline{v}, \underline{w}), y, xz, y, z
## Note we also disable the stopping stochastically as a function of size
## to allow the population to grow large and generate may different
## clones.
ospI <- oncoSimulPop(2,</pre>
                       o3init, model = "Exp",
                       mu = 5e-5, finalTime = 500,
                       detectionDrivers = 3,
                       onlyCancer = TRUE,
                       initSize = 10,
                       keepPhylog = TRUE,
                       initMutant = c("d > m > z"),
                       mc.cores = 2
                       )
op \leftarrow par(mar = rep(0, 4), mfrow = c(1, 2))
plotClonePhylog(ospI[[1]])
plotClonePhylog(ospI[[2]])
```



### 6.3 Ending the simulations

OncoSimulR provides very flexible ways to decide when to stop a simulation. Here we focus on a single simulation; see further options with multiple simulations in 7.

#### 6.3.1 Ending the simulations: conditions

- onlyCancer = TRUE. A simulation will be repeated until any one of the "reach cancer" conditions is met, if this happens before the simulation reaches finalTime<sup>11</sup>. These conditions are:
  - Total population size becomes larger than detectionSize.
  - The number of drivers in any one genotype or clone becomes equal to, or larger than, detectionDrivers; note that this allows you to stop the simulation as soon as a specific genotype is found, by using exactly and only the genes that make that genotype as the drivers. This is not allowed by the moment in frequency-dependent fitness simulations.
  - A gene or gene combination among those listed in fixation becomes fixed in the population (i.e., has a frequency is 1) (see details in (6.3.3 and 6.3.4).
  - The tumor is detected according to a stochastic detection mechanism, where the probability of "detecting the tumor" increases with population size; this is explained below (6.3.2) and is controlled by argument detectionProb.

As we exit as soon as any of the exiting conditions is reached, if you only care about one condition, set the other to NA (see also section 6.3.2.1).

• onlyCancer = FALSE. A simulation will run only once, and will exit as soon as any of the above conditions are met or as soon as the total population size becomes zero or we reach finalTime.

As an example of onlyCancer = TRUE, focusing on the first two mechanisms, suppose you give detectionSize = 1e4 and detectionDrivers =3 (and you have detectionProb = NA). A simulation will exit as soon as it reaches a total population size of 10<sup>4</sup> or any clone has four drivers, whichever comes first (if any of these happen before finalTime).

In the onlyCancer = TRUE case, what happens if we reach finalTime (or the population size becomes zero) before any of the "reach cancer" conditions have been fulfilled? The simulation will be repeated again, within the following limits:

- max.wall.time: the total wall time we allow an individual simulation to run;
- max.num.tries: the maximum number of times we allow a simulation to be repeated to reach cancer;
- max.wall.time.total and max.num.tries.total, similar to the above but over a set of simulations in function oncoSimulSample.

Incidentally, we keep track of the number of attempts used (the component other\$attemptsUsed\$) before we reach cancer, so you can estimate (as from a negative binomial sampling) the probability of reaching your desired end point under different scenarios.

<sup>&</sup>lt;sup>11</sup>Of course, the "reach cancer" idea and the onlyCancer argument are generic names; this could have been labeled "reach whatever interests me".

The onlyCancer = FALSE case might be what you want to do when you examine general population genetics scenarios without focusing on possible sampling issues. To do this, set finalTime to the value you want and set onlyCancer = FALSE; in addition, set detectionProb to "NA" and detectionDrivers and detectionSize to "NA" or to huge numbers<sup>12</sup>. In this scenario you simply collect the simulation output at the end of the run, regardless of what happened with the population (it became extinct, it did not reach a large size, it did not accumulate drivers, etc).

#### 6.3.2 Stochastic detection mechanism: "detectionProb"

This is the process that is controlled by the argument detectionProb. Here the probability of tumor detection increases with the total population size. This is biologically a reasonable assumption: the larger the tumor, the more likely it is it will be detected.

At regularly spaced times during the simulation, we compute the probability of detection as a function of size and determine (by comparing against a random uniform number) if the simulation should finish. For simplicity, and to make sure the probability is bounded between 0 and 1, we use the function

$$P(N) = \begin{cases} 1 - e^{-c((N-B)/B)} & \text{if } N > B\\ 0 & \text{if } N \le B \end{cases}$$
 (1)

where P(N) is the probability that a tumor with a population size N will be detected, and c (argument cPDetect in the oncoSimul\* functions) controls how fast P(N) increases with increasing population size relative to a baseline, B (PDBaseline in the oncoSimul\* functions); with B we both control the minimal population size at which this mechanism stats operating (because we will rarely want detection unless there is some meaningful increase of population size over initSize) and we model the increase in P(N) as a function of relative differences with respect to B. (Note that this is a major change in version 2.9.9. Before version 2.9.9, the expression used was  $P(N) = 1 - e^{-c(N-B)}$ , so we did not make the increase relative to B; of course, you can choose an appropriate c to make different models comparable, but the expression used before 2.9.9 made it much harder to compare simulations with very different initial population sizes, as baselines are often naturall a function of initial population sizes.)

The P(N) refers to the probability of detection at each one of the occasions when we assess the probability of exiting. When, or how often, do we do that? When we assess probability of exiting is controlled by checkSizePEvery, which will often be much larger than sampleEvery<sup>13</sup>. Biologically, a way to think of checkSizePEvery is "time between doctor appointments".

The setting detection Drivers and detection Size to "NA" is in fact equivalent to setting them to the largest possible numbers for these variables:  $2^{32} - 1$  and  $\infty$ , respectively.

<sup>&</sup>lt;sup>13</sup>We assess probability of exiting at every sampling time, as given by sampleEvery, that is the smallest possible sampling time that is separated from the previous time of assessment by at least checkSizePEvery. In other words, the interval between successive assessments will be the smallest multiple integer of sampleEvery that is larger than checkSizePEvery. For example, suppose

An important **warning**, though: for populations that are growing very, very fast or where some genes might have very large effects on fitness even a moderate **checkSizePEvery** of, say, 10, might be inappropriate, since populations could have increased by several orders of magnitude between successive checks. This issue is also discussed in section 2.1.1 and 2.2.

Finally, you can specify c (cPDetect) directly (you will need to set n2 and p2 to NA). However, it might be more intuitive to specify the pair n2, p2, such that P(n2) = p2 (and from that pair we solve for the value of cPDetect).

You can get a feeling for the effects of these arguments by playing with the following code, that we do not execute here for the sake of speed. Here no mutation has any effect, but there is a non-zero probability of exiting as soon as the total population size becomes larger than the initial population size. So, eventually, all simulations will exit and, as we are using the McFarland model, population size will vary slightly around the initial population size.

```
gi2 < - rep(0, 5)
names(gi2) <- letters[1:5]</pre>
oi2 <- allFitnessEffects(noIntGenes = gi2)</pre>
s5 <- oncoSimulPop(200,
                    model = "McFL",
                    initSize = 1000,
                    detectionProb = c(p2 = 0.1,
                                       n2 = 2000,
                                       PDBaseline = 1000,
                                       checkSizePEvery = 2),
                    detectionSize = NA,
                    finalTime = NA,
                    keepEvery = NA,
                    detectionDrivers = NA)
s5
hist(unlist(lapply(s5, function(x) x$FinalTime)))
```

As you decrease checkSizePEvery the distribution of "FinalTime" will resemble more and more an exponential distribution.

In this vignette, there are some further examples of using this mechanism in 6.5.4 and 6.5.2, with the default arguments.

#### 6.3.2.1 Stochastic detection mechanism and minimum number of drivers

We said above that we exit as soon as any of the conditions is reached (i.e., we use an OR operation over the exit conditions). There is a special exception to this procedure: if you set AND\_DrvProbExit = TRUE, both the number of drivers

```
sampleEvery = 2 and checkSizePEvery = 3: we will assess exiting at times 4, 8, 12, 16, \ldots If sampleEvery = 3 and checkSizePEvery = 3: we will assess exiting at times 6, 12, 18, \ldots
```

and the detectionProb mechanism condition must fulfilled. This means that the detectionProb mechanism not assessed unless the detectionDrivers condition is. Using AND\_DrvProbExit = TRUE allows to run simulations and ensure that all of the returned simulations will have at least some cells with the number of drivers as specified by detectionDrivers. Note, though, that this does not guarantee that when you sample the population, all those drivers will be detected (as this depends on the actual proportion of cells with the drivers and the settings of samplePop).

#### 6.3.3 Fixation of genes/gene combinations

In some cases we might be interested in running simulations until a particular set of genes, or gene combinations, reaches fixation. This exit condition might be more relevant than some of the above in many non-cancer-related evolutionary genetics scenarios.

Simulations will stop as soon as any of the genes or gene combinations in the vector (or list) fixation reaches a frequency of 1. These gene combinations might have non-zero intersection (i.e., they might share genes), and those genes need not be drivers. If we want simulations to only stop when fixation of those genes/gene combinations is reached, we will set all other stopping conditions to NA. It is, of course, up to you to ensure that those stopping conditions are reasonable (that they can be reached) and to use, or not, finalTime; otherwise, simulations will eventually abort (e.g., when max.wall.time or max.num.tries are reached). Since we are asking for fixation, the Exp or Bozic models will often not be appropriate here; instead, models with competition such as McFL are more appropriate.

We return here to the example from section 5.3.

Ochs and Desai explain that "Each simulated population was evolved until either the uphill genotype or valley-crossing genotype fixed." (see Ochs and Desai (2015), p.2, section "Simulations"). We will do the same here. We specify that we want to end the simulation when either the "u" or the "v, i" genotypes have reached fixation, by passing those genotype combinations as the fixation argument (in this example using fixation = c("u", "v") would have been equivalent, since the "v" genotype by itself has fitness of 0).

We want to be explicit that fixation will be the one and only condition for ending the simulations, and thus we set arguments detectionDrivers, finalTime, detectionSize and detectionProb explicitly to NA. (We set the number of repetitions only to 10 for the sake of speed when creating the vignette).

```
initS <- 20
## We use only a small number of repetitions for the sake</pre>
```

What is the frequency of each genotype among the simulations? (or, what is the frequency of fixation of each genotype?)

Note the very large variability in reaching fixation

```
head(summary(od100)[, c(1:3, 8:9)])
     NumClones TotalPopSize LargestClone FinalTime NumIter
##
              3
## 1
                           20
                                         20
                                             12080.17
                                                        483269
                                                        141425
## 2
                           27
                                         27
                                              3535.10
              4
## 3
                                                92.73
              2
                           24
                                         24
                                                          3710
## 4
              3
                           26
                                         26
                                              5055.05
                                                        202225
## 5
              3
                           21
                                         21
                                              2805.63
                                                        112238
              3
## 6
                           29
                                         29
                                               337.97
                                                         13522
```

#### 6.3.4 Fixation of genotypes

Section 6.3.3 deals with the fixation of gene/gene combinations. What if you want fixation on specific genotypes? To give an example, suppose we have a five loci genotype and suppose that you want to stop the simulations only if genotypes "A", "B, E", or "A, B, C, D, E" reach fixation. You do not want to stop it if, say, genotype "A, B, E" reaches fixation. To specify genotypes, you prepend the genotype combinations with a "\_\_,", and that tells OncoSimulR that you want fixation of **genotypes**, not just gene combinations.

An example of the differences between the mechanisms can be seen from this code:

```
## Create a simple fitness landscape
rl1 <- matrix(0, ncol = 6, nrow = 9)
colnames(rl1) <- c(LETTERS[1:5], "Fitness")</pre>
rl1[1, 6] <- 1
rl1[cbind((2:4), c(1:3))] <- 1
rl1[2, 6] <- 1.4
rl1[3, 6] <- 1.32
rl1[4, 6] <- 1.32
rl1[5, ] \leftarrow c(0, 1, 0, 0, 1, 1.5)
rl1[6, ] \leftarrow c(0, 0, 1, 1, 0, 1.54)
rl1[7, ] <- c(1, 0, 1, 1, 0, 1.65)
rl1[8, ] <- c(1, 1, 1, 1, 0, 1.75)
rl1[9, ] \leftarrow c(1, 1, 1, 1, 1, 1.85)
class(rl1) <- c("matrix", "genotype_fitness_matrix")</pre>
## plot(rl1) ## to see the fitness landscape
## Gene combinations
local_max_g <- c("A", "B, E", "A, B, C, D, E")
## Specify the genotypes
local_max <- paste0("_,", local_max_g)</pre>
fr1 <- allFitnessEffects(genotFitness = rl1, drvNames = LETTERS[1:5])</pre>
initS <- 2000
####### Stop on gene combinations
                                        #####
r1 <- oncoSimulPop(10,
                        fp = fr1,
                        model = "McFL",
                        initSize = initS,
                        mu = 1e-4,
                        detectionSize = NA,
                        sampleEvery = .03,
                        keepEvery = 1,
                        finalTime = 50000,
                        fixation = local_max_g,
                        detectionDrivers = NA,
                        detectionProb = NA,
                        onlyCancer = TRUE,
                        max.num.tries = 500,
                        max.wall.time = 20,
                        errorHitMaxTries = TRUE,
                        keepPhylog = FALSE,
                        mc.cores = 2)
sp1 <- samplePop(r1, "last", "singleCell")</pre>
##
```

```
## Subjects by Genes matrix of 10 subjects and 5 genes.
sgsp1 <- sampledGenotypes(sp1)</pre>
## often you will stop on gene combinations that
## are not local maxima in the fitness landscape
sgsp1
##
       Genotype Freq
## 1
              \boldsymbol{A}
## 2 A, B, C, D
     A, C, D
## 3
## 4
          B, E
                    3
##
## Shannon's diversity (entropy) of sampled genotypes: 1.168
sgsp1$Genotype %in% local max g
## [1] TRUE FALSE FALSE TRUE
###### Stop on genotypes
                             ####
r2 <- oncoSimulPop(10,
                       fp = fr1,
                       model = "McFL",
                       initSize = initS,
                       mu = 1e-4.
                       detectionSize = NA,
                       sampleEvery = .03,
                       keepEvery = 1,
                       finalTime = 50000,
                       fixation = local max,
                       detectionDrivers = NA,
                       detectionProb = NA,
                        onlyCancer = TRUE,
                       max.num.tries = 500,
                       max.wall.time = 20,
                        errorHitMaxTries = TRUE,
                       keepPhylog = FALSE,
                       mc.cores = 2)
## All final genotypes should be local maxima
sp2 <- samplePop(r2, "last", "singleCell")</pre>
##
## Subjects by Genes matrix of 10 subjects and 5 genes.
sgsp2 <- sampledGenotypes(sp2)</pre>
sgsp2$Genotype %in% local max g
## [1] TRUE TRUE TRUE
```

#### 6.3.5 Fixation: tolerance, number of periods, minimal size

In particular if you specify stopping on genotypes, you might want to think about three additional parameters: fixation\_tolerance, min\_successive\_fixation, and fixation\_min\_size.

fixation\_tolerance: fixation is considered to have happened if the genotype/gene combinations specified as genotypes/gene combinations for fixation have reached a frequency  $> 1 - fixation\_tolerance$ . (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.)

min\_successive\_fixation: 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).

fixation\_min\_size: 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).

An example of using those options:

```
## Create a simple fitness landscape
rl1 \leftarrow matrix(0, ncol = 6, nrow = 9)
colnames(rl1) <- c(LETTERS[1:5], "Fitness")</pre>
rl1[1, 6] <- 1
rl1[cbind((2:4), c(1:3))] <- 1
rl1[2, 6] <- 1.4
rl1[3, 6] <- 1.32
rl1[4, 6] <- 1.32
rl1[5, ] \leftarrow c(0, 1, 0, 0, 1, 1.5)
rl1[6, ] \leftarrow c(0, 0, 1, 1, 0, 1.54)
rl1[7, ] \leftarrow c(1, 0, 1, 1, 0, 1.65)
rl1[8, ] <- c(1, 1, 1, 1, 0, 1.75)
rl1[9, ] <- c(1, 1, 1, 1, 1, 1.85)
class(rl1) <- c("matrix", "genotype fitness matrix")</pre>
## plot(rl1) ## to see the fitness landscape
## The local fitness maxima are
## c("A", "B, E", "A, B, C, D, E")
fr1 <- allFitnessEffects(genotFitness = rl1, drvNames = LETTERS[1:5])</pre>
initS <- 2000
## Stop on genotypes
```

```
r3 <- oncoSimulPop(10,
                  fp = fr1,
                  model = "McFL",
                  initSize = initS,
                  mu = 1e-4,
                  detectionSize = NA,
                  sampleEvery = .03,
                  keepEvery = 1,
                  finalTime = 50000,
                  fixation = c(paste0(" ,",
                                    c("A", "B, E", "A, B, C, D, E")),
                                fixation tolerance = 0.1,
                                min_successive_fixation = 200,
                                fixation min size = 3000),
                  detectionDrivers = NA,
                  detectionProb = NA,
                  onlyCancer = TRUE,
                  max.num.tries = 500,
                  max.wall.time = 20,
                  errorHitMaxTries = TRUE,
                  keepPhylog = FALSE,
                  mc.cores = 2)
```

#### 6.3.6 Mixing stopping on gene combinations and genotypes

This would probably be awfully confusing and is not tested formally (though it should work). Let me know if you think this is an important feature. (Pull requests with tests welcome.)

## 6.4 Plotting genotype/driver abundance over time; plotting the simulated trajectories

We have seen many of these plots already, starting with Figure 2 and Figure 4 and we will see many more below, in the examples, starting with section 6.5.1 such as in figures 5 and 6. In a nutshell, what we are plotting is the information contained in the pops.by.time matrix, the matrix that contains the abundances of all the clones (or genotypes) at each of the sampling periods.

The functions that do the work are called plot and these are actually methods for objects of class "oncosimul" and "oncosimulpop". You can access the help by doing <code>?plot.oncosimul</code>, for example.

What entities are shown in the plot? You can show the trajectories of:

- numbers of drivers (e.g., 5);
- genotypes or clones (e.g., 6).

(Of course, showing "drivers" requires that you have specified certain genes as drivers.)

What types of plots are available?

- line plots;
- stacked plots;
- stream plots.

All those three are shown in both of Figure 5 and Figure 6.

If you run multiple simulations using oncoSimulPop you can plot the trajectories of all of the simulations.

## 6.5 Several examples of simulations and plotting simulation trajectories

#### 6.5.1 Bauer's example again

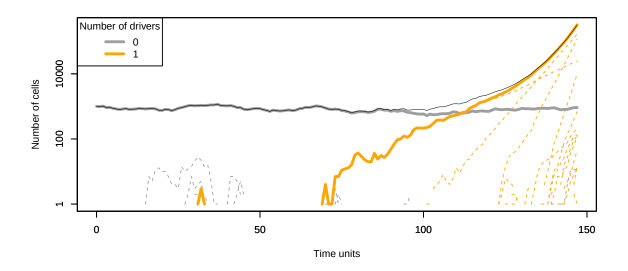
We will use the model of Bauer, Siebert, and Traulsen (2014) that we saw in section 5.1.

```
K <- 5
sd <- 0.1
sdp <- 0.15
sp < -0.05
bauer <- data.frame(parent = c("Root", rep("p", K)),</pre>
                     child = c("p", paste0("s", 1:K)),
                     s = c(sd, rep(sdp, K)),
                     sh = c(0, rep(sp, K)),
                     typeDep = "MN")
fbauer <- allFitnessEffects(bauer, drvNames = "p")</pre>
set.seed(1)
## Use fairly large mutation rate
b1 <- oncoSimulIndiv(fbauer, mu = 5e-5, initSize = 1000,
                      finalTime = NA,
                      onlyCancer = TRUE,
                      detectionProb = "default")
```

We will now use a variety of plots

```
par(mfrow = c(3, 1))
## First, drivers
plot(b1, type = "line", addtot = TRUE)
plot(b1, type = "stacked")
plot(b1, type = "stream")

par(mfrow = c(3, 1))
## Next, genotypes
plot(b1, show = "genotypes", type = "line")
```



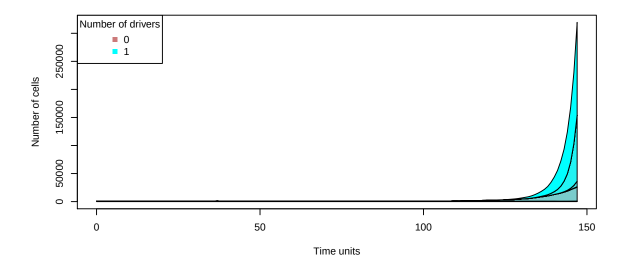




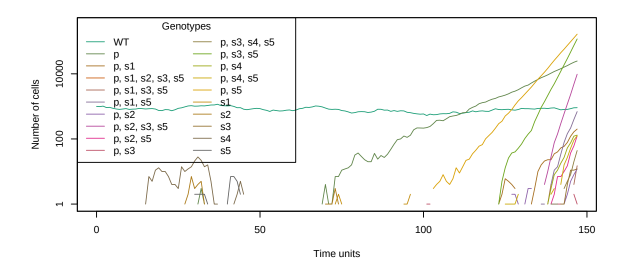
Figure 5: Three drivers' plots of a simulation of Bauer's model

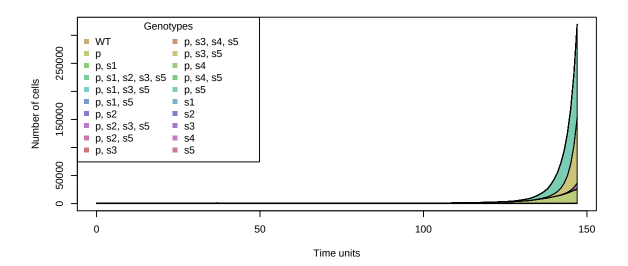
```
plot(b1, show = "genotypes", type = "stacked")
plot(b1, show = "genotypes", type = "stream")
```

In this case, probably the stream plots are most helpful. Note, however, that (in contrast to some figures in the literature showing models of clonal expansion) the stream plot (or the stacked plot) does not try to explicitly show parent-descendant relationships, which would hardly be realistically possible in these plots (although the plots of phylogenies in section 8 could be of help).

#### 6.5.2 McFarland model with 5000 passengers and 70 drivers

```
set.seed(678)
nd <- 70
np <- 5000
s < -0.1
sp <- 1e-3
spp <- -sp/(1 + sp)
mcf1 <- allFitnessEffects(noIntGenes = c(rep(s, nd), rep(spp, np)),</pre>
                           drvNames = seq.int(nd))
mcf1s <- oncoSimulIndiv(mcf1,</pre>
                          model = "McFL",
                          mu = 1e-7,
                          detectionProb = "default",
                          detectionSize = NA,
                          detectionDrivers = NA,
                          sampleEvery = 0.025,
                          keepEvery = 8,
                          initSize = 2000,
                          finalTime = 4000,
                          onlyCancer = FALSE)
summary(mcf1s)
##
     {\it NumClones} {\it TotalPopSize} {\it LargestClone} {\it MaxNumDrivers} {\it MaxDriversLast}
                                                         3
## 1
            618
                         3207
                                       2841
                                                                          3
##
     NumDriversLargestPop TotalPresentDrivers FinalTime NumIter
## 1
                          3
                                                5
                                                       1961
##
     HittedWallTime HittedMaxTries errorMF minDMratio minBMratio
## 1
               FALSE
                               FALSE 0.0129
                                                     1585
                                                                 1972
##
      OccurringDrivers
## 1 5, 24, 30, 57, 67
par(mfrow = c(2, 1))
## I use thinData to make figures smaller and faster
plot(mcf1s, addtot = TRUE, lwdClone = 0.9, log = "",
     thinData = TRUE, thinData.keep = 0.5)
plot(mcf1s, show = "drivers", type = "stacked",
     thinData = TRUE, thinData.keep = 0.3,
```





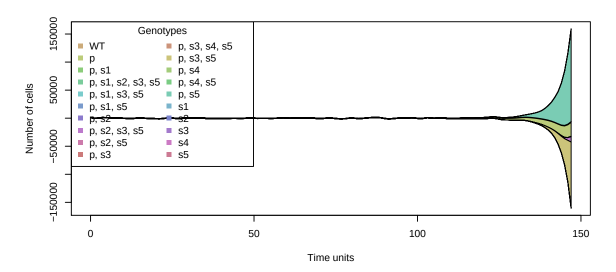
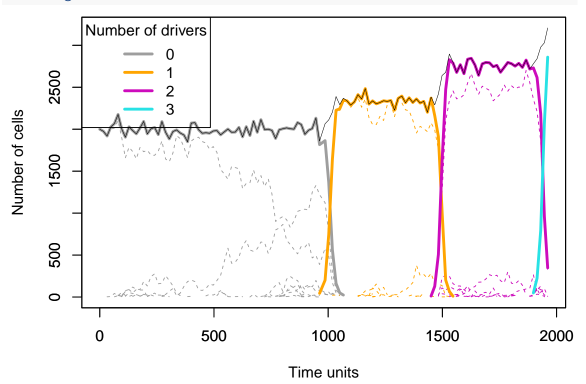
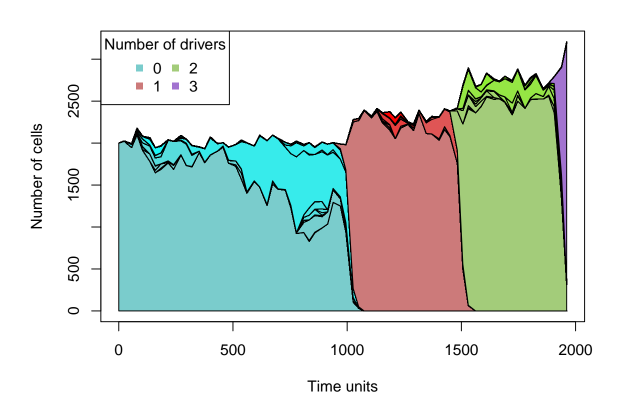


Figure 6: Three genotypes' plots of a simulation of Bauer's model







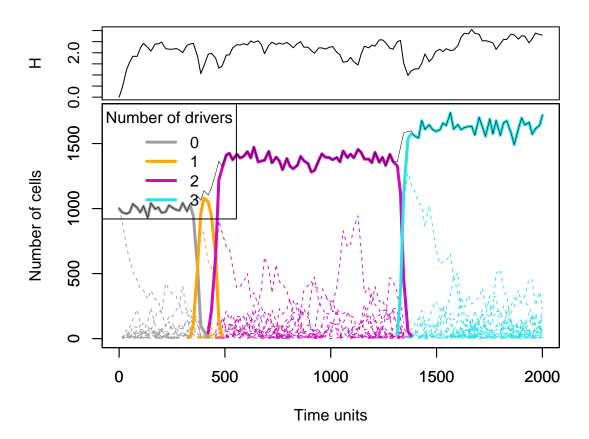
With the above output (where we see there are over 500 different genotypes) trying to represent the genotypes makes no sense.

## 6.5.3 McFarland model with 50,000 passengers and 70 drivers: clonal competition

The next is too slow (takes a couple of minutes in an i5 laptop) and too big to run in a vignette, because we keep track of over 4000 different clones (which leads to a result object of over 800 MB):

```
set.seed(123)
nd <- 70
np <- 50000
s < -0.1
sp <- 1e-4 ## as we have many more passengers
spp < - -sp/(1 + sp)
mcfL <- allFitnessEffects(noIntGenes = c(rep(s, nd), rep(spp, np)),</pre>
                           drvNames = seq.int(nd))
mcfLs <- oncoSimulIndiv(mcfL,</pre>
                          model = "McFL",
                          mu = 1e-7,
                          detectionSize = 1e8,
                          detectionDrivers = 100,
                          sampleEvery = 0.02,
                          keepEvery = 2,
                          initSize = 1000,
                          finalTime = 2000,
                          onlyCancer = FALSE)
```

But you can access the pre-stored results and plot them (beware: this object has been trimmed by removing empty passenger rows in the Genotype matrix)

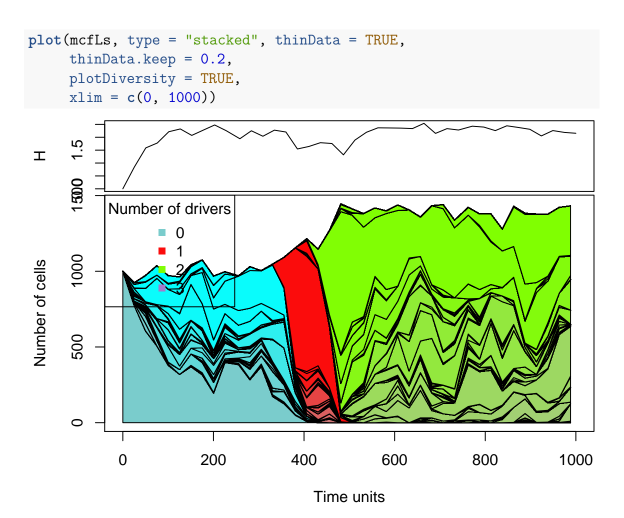


The argument plotDiversity = TRUE asks to show a small plot on top with Shannon's diversity index.

```
summary(mcfLs)
##
     NumClones TotalPopSize LargestClone MaxNumDrivers MaxDriversLast
## 1
          4458
                        1718
                                       253
     {\it NumDriversLargestPop\ TotalPresentDrivers\ FinalTime\ NumIter}
##
## 1
                                              70
                                                      2000
                                                             113759
##
     HittedWallTime HittedMaxTries errorMF minDMratio minBMratio
## 1
              FALSE
                              FALSE 0.01922
                                                   184.1
                                                               199.6
     OccurringDrivers
##
       13, 38, 40, 69
## number of passengers per clone
summary(colSums(mcfLs$Genotypes[-(1:70), ]))
##
      Min. 1st Qu.
                     Median
                               Mean 3rd Qu.
                                                 Max.
##
      0.00
              4.00
                       6.00
                                5.67
                                        7.75
                                                13.00
```

Note that we see clonal competition between clones with the same number of drivers (and with different drivers, of course). We will return to this (section 6.5.5).

A stacked plot might be better to show the extent of clonal competition (plotting takes some time —a stream plot reveals similar patterns and is also slower than the line plot). I will aggressively thin the data for this plot so it is faster and smaller (but we miss some of the fine grain, of course):



#### 6.5.4 Simulation with a conjunction example

We will use several of the previous examples. Most of them are in file examplesFitnessEffects, where they are stored inside a list, with named components (names the same as in the examples above):

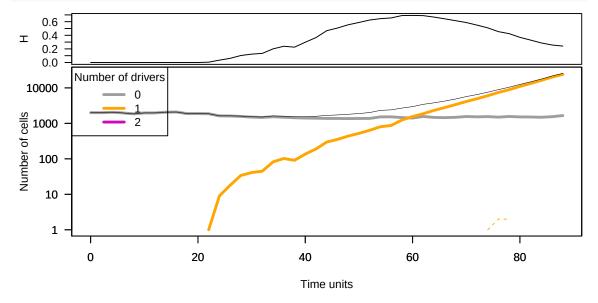
```
data(examplesFitnessEffects)
names(examplesFitnessEffects)
    [1] "cbn1"
                  "cbn2"
                                      "xor1"
                                                                   "03"
##
                            "smn1"
                                                "fp3"
                                                          "fp4m"
##
    [8] "ofe1"
                  "ofe2"
                            "foi1"
                                      "sv"
                                                "svB"
                                                         "svB1"
                                                                   "sv2"
                                                         "fbauer" "w"
                  "e2"
                            "E3A"
## [15] "sm1"
                                      "em"
                                                "fea"
## [22] "pancr"
```

We will simulate using the simple CBN-like restrictions of section 3.4.4 with two different models.

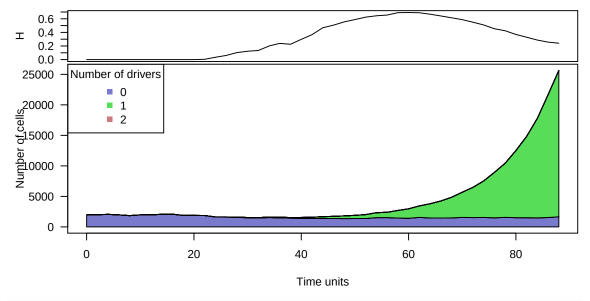
```
## 3
             c 0.10
## 4
             d
                  1.10
## 5
                 1.10
             e
## 6
                 0.10
             g
          a, b
## 7
                 1.21
## 8
                 0.11
          a, c
## 9
          a, d
                 1.21
## 10
                  1.21
          a, e
sm <- oncoSimulIndiv(examplesFitnessEffects$cbn1,</pre>
                      model = "McFL",
                      mu = 5e-7,
                      detectionSize = 1e8,
                      detectionDrivers = 2,
                      detectionProb = "default",
                      sampleEvery = 0.025,
                      keepEvery = 5,
                      initSize = 2000,
                      onlyCancer = TRUE)
summary(sm)
     NumClones TotalPopSize LargestClone MaxNumDrivers MaxDriversLast
             6
                       2721
                                                      2
## 1
                                     2005
##
     NumDriversLargestPop TotalPresentDrivers FinalTime NumIter
                                             4
   HittedWallTime HittedMaxTries errorMF minDMratio minBMratio
## 1
              FALSE
                            FALSE 0.01502
                                                309424
                                                           333333
    OccurringDrivers
##
## 1
           a, b, c, e
set.seed(1234)
evalAllGenotypes(examplesFitnessEffects$cbn1, order = FALSE,
                 model = "Bozic")[1:10, ]
##
      Genotype Death_rate
## 1
                     0.90
             \boldsymbol{a}
## 2
                     0.90
## 3
                     1.90
            C
## 4
            d
                     0.90
## 5
                     0.90
            е
## 6
                     1.90
             g
## 7
                     0.81
          a, b
                     1.71
## 8
          a, c
## 9
          a, d
                     0.81
                     0.81
## 10
          a, e
sb <- oncoSimulIndiv(examplesFitnessEffects$cbn1,</pre>
                       model = "Bozic",
                       mu = 5e-6,
                      detectionProb = "default",
```

```
detectionSize = 1e8,
                       detectionDrivers = 4,
                       sampleEvery = 2,
                       initSize = 2000,
                       onlyCancer = TRUE)
summary(sb)
##
     NumClones TotalPopSize LargestClone MaxNumDrivers MaxDriversLast
                       25636
                                    23975
## 1
##
     NumDriversLargestPop TotalPresentDrivers FinalTime NumIter
## 1
                                             5
                                                       88
##
    HittedWallTime HittedMaxTries errorMF minDMratio minBMratio
## 1
              FALSE
                              FALSE
                                         NA
                                                  33333
                                                             33333
##
     OccurringDrivers
        a, b, c, e, g
```

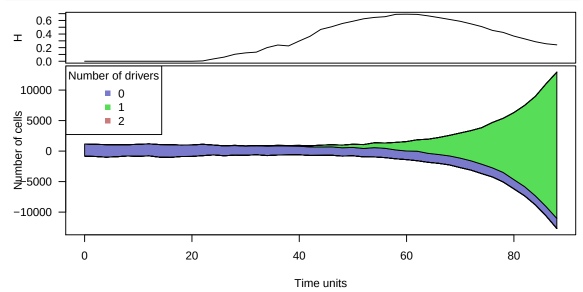
As usual, we will use several plots here.



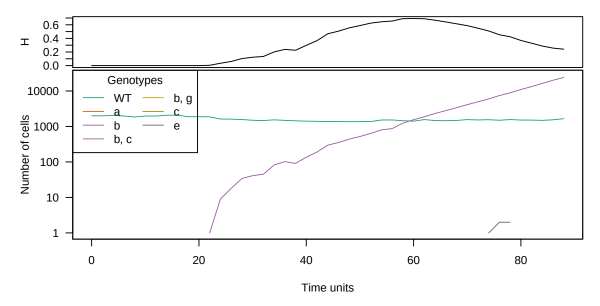
```
## Drivers, stacked
par(cex = 0.75, las = 1)
plot(sb,show = "drivers", type = "stacked", plotDiversity = TRUE)
```



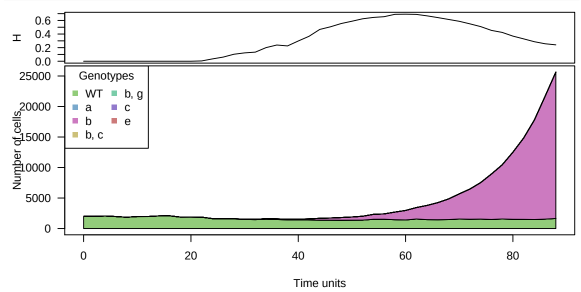
```
## Drivers, stream
par(cex = 0.75, las = 1)
plot(sb,show = "drivers", type = "stream", plotDiversity = TRUE)
```



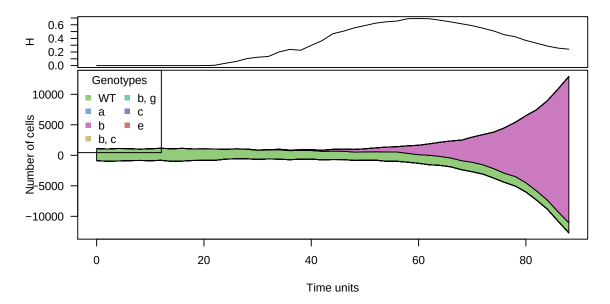
```
## Genotypes, line plot
par(cex = 0.75, las = 1)
plot(sb,show = "genotypes", type = "line", plotDiversity = TRUE)
```



```
## Genotypes, stacked
par(cex = 0.75, las = 1)
plot(sb,show = "genotypes", type = "stacked", plotDiversity = TRUE)
```



```
## Genotypes, stream
par(cex = 0.75, las = 1)
plot(sb,show = "genotypes", type = "stream", plotDiversity = TRUE)
```

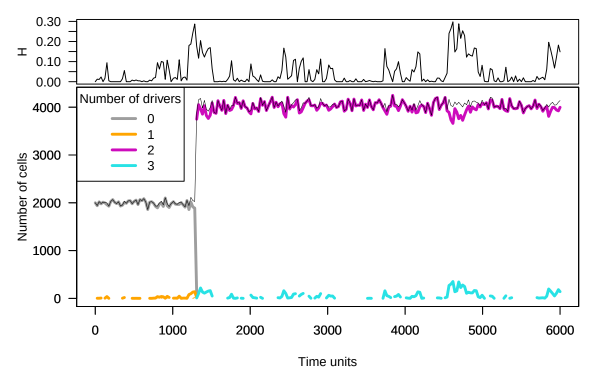


The above illustrates again that different types of plots can be useful to reveal different patterns in the data. For instance, here, because of the huge relative frequency of one of the clones/genotypes, the stacked and stream plots do not reveal the other clones/genotypes as we cannot use a log-transformed y-axis, even if there are other clones/genotypes present.

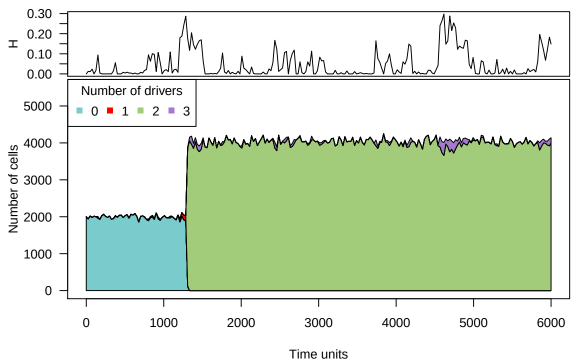
#### 6.5.5 Simulation with order effects and McFL model

(We use a somewhat large mutation rate than usual, so that the simulation runs quickly.)

We show a stacked and a line plot of the drivers:



```
par(las = 1, cex = 0.85)
plot(tmp, type = "stacked", plotDiversity = TRUE,
    ylim = c(0, 5500), legend.ncols = 4,
    thinData = TRUE, thinData.keep = 0.2)
```

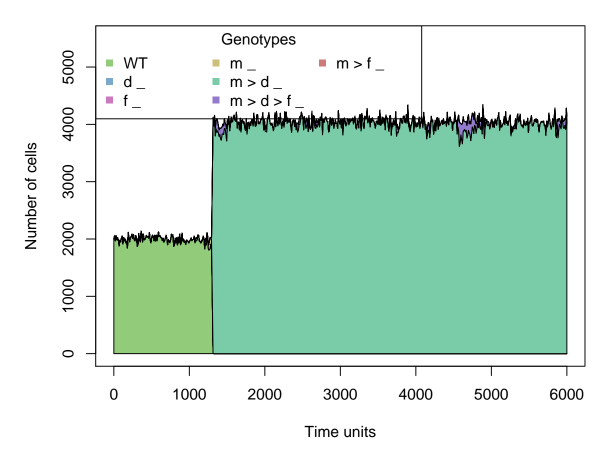


In this example (and at least under Linux, with both GCC and clang —random number streams in C++, and thus simulations, can differ between combinations of operating system and compiler), we can see that the mutants with three drivers do

not get established when we stop the simulation at time 6000. This is one case where the summary statistics about number of drivers says little of value, as fitness is very different for genotypes with the same number of mutations, and does not increase in a simple way with drivers:

```
evalAllGenotypes(examplesFitnessEffects[["o3"]], addwt = TRUE,
                 order = TRUE)
##
       Genotype Fitness
## 1
             WT
                   1.00
## 2
              d
                   1.00
## 3
              f
                   1.00
## 4
                   1.00
              m
## 5
          d > f
                   1.00
## 6
          d > m
                   1.10
          f > d
## 7
                   1.00
## 8
          f > m
                   1.00
## 9
          m > d
                   1.50
## 10
          m > f
                   1.00
## 11 d > f > m
                   1.54
## 12 d > m > f
                   1.32
                   0.77
## 13 f > d > m
## 14 f > m > d
                   1.50
## 15 m > d > f
                   1.50
## 16 m > f > d
                   1.50
```

A few figures could help:

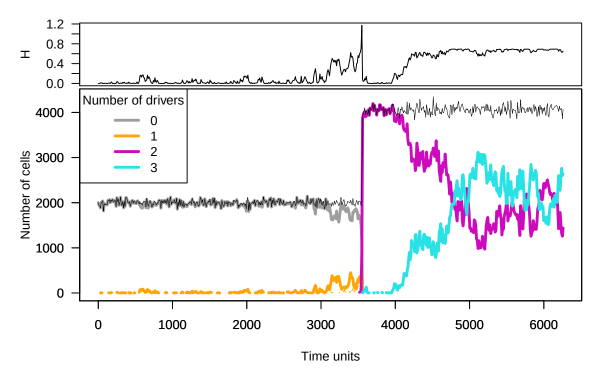


(When reading the figure legends, recall that genotype  $x > y \_ z$  is one where a mutation in "x" happened before a mutation in "y", and there is also a mutation in "z" for which order does not matter. Here, there are no genes for which order does not matter and thus there is nothing after the " $\_$ ").

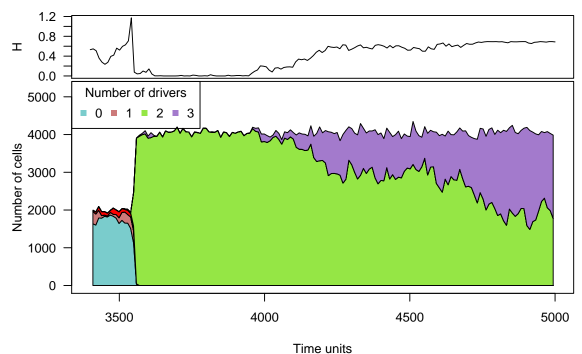
In this case, the clones with three drivers end up displacing those with two by the time we stop; moreover, notice how those with one driver never really grow to a large population size, so we basically go from a population with clones with zero drivers to a population made of clones with two or three drivers:

```
## Individual OncoSimul trajectory with call:
   oncoSimulIndiv(fp = examplesFitnessEffects[["o3"]], model = "McFL",
      mu = 5e-05, detectionSize = 1e+08, detectionDrivers = 3,
      sampleEvery = 0.025, initSize = 2000, keepEvery = 5, extraTime = 1500,
##
      finalTime = 20000, onlyCancer = FALSE, max.num.tries = 10)
##
##
##
    \it NumClones \ TotalPopSize \ LargestClone \ \it MaxNumDrivers \ \it MaxDriversLast
                                   2632
## 1
                      4071
    {\it NumDriversLargestPop\ TotalPresentDrivers\ FinalTime\ NumIter}
##
## 1
                       3
                                           3
                                                 6259 251954
##
   HittedWallTime HittedMaxTries errorMF minDMratio minBMratio
## 1
             FALSE
                           FALSE 0.01442
                                             6161
                                                           6667
## OccurringDrivers
## 1
             d, f, m
##
## Final population composition:
       Genotype
                   N
## 1
                     0
## 2
                    0
           d \_
## 3
       d > f
                    0
## 4
            f_{-}
## 5
##6  m>d
                1439
## 7 m > d > f
                2632
## 8
      m > f
```

use a drivers plot:

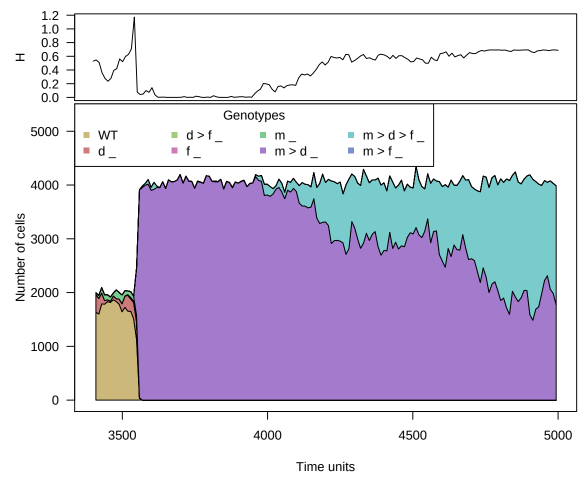


```
par(las = 1, cex = 0.85)
plot(tmp, type = "stacked", plotDiversity = TRUE,
    legend.ncols = 4, ylim = c(0, 5200), xlim = c(3400, 5000),
    thinData = TRUE, thinData.keep = 0.5)
```



Now show the genotypes explicitly:

```
## Improve telling apart the most abundant
## genotypes by sorting colors
## differently via breakSortColors
```



As before, the argument plotDiversity = TRUE asks to show a small plot on top with Shannon's diversity index. Here, as before, the quick clonal expansion of the clone with two drivers leads to a sudden drop in diversity (for a while, the population is made virtually of a single clone). Note, however, that compared to section 6.5.3, we are modeling here a scenario with very few genes, and correspondingly very few possible genotypes, and thus it is not strange that we observe very little diversity.

(We have used extraTime to continue the simulation well past the point of detection, here specified as three drivers. Instead of specifying extraTime we can set the detectionDrivers value to a number larger than the number of existing possible drivers, and the simulation will run until finalTime if onlyCancer = FALSE.)

### 6.6 Interactive graphics

It is possible to create interactive stacked area and stream plots using the *streamgraph* package, available from https://github.com/hrbrmstr/streamgraph. However, that package is not available as a CRAN or BioConductor package, and thus we cannot depend on it for this vignette (or this package). You can, however, paste the code below and make it run locally.

Before calling the streamgraph function, though, we need to convert the data from the original format in which it is stored into "long format". A simple convenience function is provided as OncoSimulWide2Long in OncoSimulR.

As an example, we will use the data we generated above for section 6.5.1.

## 7 Sampling multiple simulations

Often, you will want to simulate multiple runs of the same scenario, and then obtain the matrix of runs by mutations (a matrix of individuals/samples by genes or, equivalently, a vector of "genotypes"), and do something with them. OncoSimulR offers several ways of doing this.

The key function here is samplePop, either called explicitly after oncoSimulPop (or oncoSimulIndiv), or implicitly as part of a call to oncoSimulSample. With samplePop you can use single cell or whole tumor sampling (for details see the help of samplePop). Depending on how the simulations were conducted, you might also sample at different times, or as a function of population sizes. A major difference between procedures has to do with whether or not you want to keep the complete history of the simulations.

You want to keep the complete history of population sizes of clones during the simulations. You will simulate using:

- oncoSimulIndiv repeatedly (maybe within mclapply, to parallelize the run).
- oncoSimulPop. oncoSimulPop is basically a thin wrapper around oncoSimulIndiv that uses mclapply.

In both cases, you specify the conditions for ending the simulations (as explained in 6.3). Then, you use function samplePop to obtain the matrix of samples by mutations.

You do not want to keep the complete history of population sizes of clones during the simulations. You will simulate using:

- oncoSimulIndiv repeatedly, with argument keepEvery = NA.
- oncoSimulPop, with argument keepEvery = NA.
  - In both cases you specify the conditions for ending the simulations (as explained in 6.3). Then, you use function samplePop.
- oncoSimulSample, specifying the conditions for ending the simulations (as explained in 6.3). In this case, you will not use samplePop, as that is implicitly called by oncoSimulSample. The output is directly the matrix (and a little bit of summary from each run), and during the simulation it only stores one time point.

Why the difference between the above cases? If you keep the complete history of population sizes, you can take samples at any of the times between the beginning and the end of the simulations. If you do not keep the history, you can only sample at the time the simulation exited (see section 14.2). Why would you want to use the second route? If we are only interested in the final matrix of individuals by mutations, keeping the complete history above is wasteful because we store fully all of the simulations (for example in the call to oncoSimulPop) and then sample (in the call to samplePop). Further criteria to use when choosing between sampling procedures is whether you need detectionSize and detectionDrivers do differ

between simulations: if you use oncoSimulPop the arguments for detectionSize and detectionDrivers must be the same for all simulations but this is not the case for oncoSimulSample. See further comments in 7.2. Finally, parallelized execution is available for oncoSimulPop but, by design, not for oncoSimulSample.

The following are a few examples. First we run oncoSimulPop to obtain 4 simulations and in the last line we sample from them:

```
pancrPop <- oncoSimulPop(4, pancr,</pre>
                           detectionSize = 1e7,
                           keepEvery = 10,
                           mc.cores = 2)
summary(pancrPop)
##
     NumClones TotalPopSize LargestClone MaxNumDrivers MaxDriversLast
## 1
             12
                     10603644
                                   10229581
                                                           0
                                                                           0
                                                           0
                                                                           0
## 2
             14
                     10987604
                                    7122102
## 3
                                                           0
                                                                           0
             18
                     11473182
                                    5893155
## 4
             10
                     10877452
                                   10841003
                                                           0
                                                                           0
##
     NumDriversLargestPop TotalPresentDrivers FinalTime NumIter
## 1
                          0
                                                 0
                                                        1264
                                                                 1919
                          0
                                                 0
## 2
                                                         912
                                                                 1496
## 3
                          0
                                                 0
                                                        2047
                                                                 2610
## 4
                          0
                                                        1049
                                                                 1827
     HittedWallTime HittedMaxTries errorMF minDMratio minBMratio
##
## 1
               FALSE
                                FALSE
                                            NA
                                                    142857
                                                                142857
## 2
               FALSE
                                FALSE
                                                    142857
                                                                142857
                                            NA
## 3
               FALSE
                                FALSE
                                            NA
                                                    142857
                                                                142857
               FALSE
                                FALSE
                                                    142857
## 4
                                            NA
                                                                142857
     OccurringDrivers
## 1
## 2
## 3
## 4
samplePop(pancrPop)
##
##
    Subjects by Genes matrix of 4 subjects and 7 genes.
         CDNK2A KRAS MLL3 PXDN SMAD4 TGFBR2 TP53
##
## [1,]
              0
                    1
                         0
                               0
                                      0
                                             0
## [2,]
              0
                    1
                         0
                               0
                                      0
                                             0
                                                   0
## [3,]
              0
                    1
                         0
                               0
                                      0
                                             0
                                                   0
                               0
                                      0
              0
                    1
                                                   0
## [4,]
```

Now a simple multiple call to oncoSimulIndiv wrapped inside mclapply; this is basically the same we just did above. We set the class of the object to allow direct usage of samplePop. (Note: in Windows mc.cores > 1 is not supported, so for the vignette to run in Windows, Linux, and Mac we explicitly set it here in the call

to mclapply. For regular usage, you will not need to do this; just use whatever is appropriate for your operating system and number of cores. As well, we do not need any of this with oncoSimulPop because the code inside oncoSimulPop already takes care of setting mc.cores to 1 in Windows).

```
library(parallel)
if(.Platform$0S.type == "windows") {
    mc.cores <- 1
} else {
    mc.cores <- 2
}
p2 <- mclapply(1:4, function(x) oncoSimulIndiv(pancr,
                                                  detectionSize = 1e7,
                                                  keepEvery = 10),
                                                  mc.cores = mc.cores)
class(p2) <- "oncosimulpop"</pre>
samplePop(p2)
##
##
    Subjects by Genes matrix of 4 subjects and 7 genes.
##
        CDNK2A KRAS MLL3 PXDN SMAD4 TGFBR2 TP53
## [1,]
              0
                    1
                         0
                               0
                                     0
                                             0
                                                  0
## [2,]
                         0
                               0
                                     0
                                             0
                                                   0
              0
                    1
## [3,]
              0
                    1
                         0
                               0
                                     0
                                             0
                                                   0
                    1
                         0
                                     0
                                                   1
## [4,]
              0
```

Above, we have kept the complete history of the simulations as you can check by doing, for instance

```
tail(pancrPop[[1]]$pops.by.time)
##
           [,1] [,2]
                       [,3] [,4] [,5]
                                            [,6] [,7] [,8] [,9] [,10]
## [123,] 1220 2128
                         36
                                0
                                          125991
                                                    0
                                                          0
                                                               0
                                                                      5
## [124,] 1230 2200
                        288
                                0
                                     0
                                          342171
                                                    0
                                                          0
                                                               0
                                                                     13
                                0
                                                               2
## [125,] 1240 1806
                       2296
                                     0
                                          928133
                                                    1
                                                          0
                                                                    132
                                                               5
## [126,] 1250 1813
                      18559
                                0
                                     0 2523665
                                                   29
                                                          0
                                                                   1231
## [127,] 1260 1788 150079
                                9
                                                          5
                                                                  9785
                                     0
                                        6860270
                                                  240
                                                               3
## [128,] 1264 1764 344967
                               15
                                     1 10229581
                                                  528
                                                         31
                                                              14 22581
          [,11] [,12] [,13]
##
## [123,]
               0
                     0
                            0
## [124,]
               0
                     0
                            0
                            0
## [125,]
               0
                    14
## [126,]
               2
                   247
                            0
## [127,]
               3
                  1792
                            0
                            0
## [128,]
               8
                  4154
```

If we were not interested in the complete history of simulations we could have done instead (note the argument keepEvery = NA)

```
pancrPopNH <- oncoSimulPop(4, pancr,</pre>
                            detectionSize = 1e7,
                            keepEvery = NA,
                            mc.cores = 2)
summary(pancrPopNH)
     NumClones TotalPopSize LargestClone MaxNumDrivers MaxDriversLast
## 1
                    10343468
                                  10335506
                                                        0
                                                                        0
             8
## 2
             8
                    10733600
                                  10567729
                                                        0
                                                                        0
## 3
             15
                    10253954
                                   8343771
                                                        0
                                                                        0
## 4
             10
                    10601365
                                  10589985
                                                        0
                                                                        0
     NumDriversLargestPop TotalPresentDrivers FinalTime NumIter
## 1
                         0
                                               0
                                                       805
                                                               1501
                         0
## 2
                                               0
                                                       225
                                                                906
## 3
                         0
                                               0
                                                       995
                                                               1646
                         0
                                               0
                                                      1966
                                                               2677
## 4
## HittedWallTime HittedMaxTries errorMF minDMratio minBMratio
## 1
              FALSE
                              FALSE
                                          NA
                                                  142857
                                                             142857
## 2
              FALSE
                              FALSE
                                          NA
                                                  142857
                                                              142857
## 3
              FALSE
                              FALSE
                                          NA
                                                  142857
                                                             142857
## 4
              FALSE
                              FALSE
                                          NA
                                                  142857
                                                             142857
##
     OccurringDrivers
## 1
## 2
## 3
## 4
samplePop(pancrPopNH)
##
##
   Subjects by Genes matrix of 4 subjects and 7 genes.
        CDNK2A KRAS MLL3 PXDN SMAD4 TGFBR2 TP53
##
## [1,]
             0
                   1
                        0
                             0
                                    0
                                            0
## [2,]
                        0
                              0
                                    0
                                            0
                                                 0
              0
                   1
## [3,]
                                    0
                                            0
                                                 0
              0
                   1
                        0
                              0
## [4,]
              0
                   1
```

which only keeps the very last sample:

```
pancrPopNH[[1]]$pops.by.time
## [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9]
## [1,] 805 418 2273 10335506 685 18 1187 20 3361
```

Or we could have used oncoSimulSample:

```
pancrSamp <- oncoSimulSample(4, pancr)
## Successfully sampled 4 individuals
##
## Subjects by Genes matrix of 4 subjects and 7 genes.</pre>
```

pan	pancrSamp <mark>\$</mark> popSamp							
##		CDNK2A	KRAS	MLL3	PXDN	SMAD4	TGFBR2	<i>TP53</i>
##	[1,]	0	1	0	0	0	0	0
##	[2,]	0	1	0	0	0	0	0
##	[3,]	0	1	0	0	0	0	0
##	[4,]	0	1	0	0	0	0	0

Again, why the above differences? If we are only interested in the final matrix of populations by mutations, keeping the complete history the above is wasteful, because we store fully all of the simulations (in the call to oncoSimulPop) and then sample (in the call to samplePop).

# 7.1 Whole-tumor and single-cell sampling, and do we always want to sample?

samplePop is designed to emulate the process of obtaining a sample from a (set of) "patient(s)". But there is no need to sample. The history of the population, with a granularity that is controlled by argument keepEvery, is kept in the matrix pops.by.time which contains the number of cells of every clone at every sampling point (see further details in 14.2). This is the information used in the plots that show the trajectory of a simulation: the plots that show the change in genotype or driver abundance over time (see section 6.4 and examples mentioned there).

Regardless of whether and how you plot the information in pops.by.time, you can also sample one or multiple simulations using samplePop. In whole-tumor sampling the resolution is the whole tumor (or the whole population). Thus, a key argument is thresholdWhole, the threshold for detecting a mutation: a gene is considered mutated if it is altered in at least "thresholdWhole" proportion of the cells in that simulation (at a particular time point). This of course means that your "sampled genotype" might not correspond to any existing genotype because we are summing over all cells in the population. For instance, suppose that at the time we take the sample there are only two clones in the population, one clone with a frequency of 0.4 that has gene A mutated, and a second clone one with a frequency of 0.6 that has gene B mutated. If you set thresholdWhole to values  $\leq 0.4$  the sampled genotype will show both A and B mutated. Single-cell sampling is provided as an option in contrast to whole-tumor sampling. Here any sampled genotype will correspond to an existing genotype as you are sampling with single-cell resolution.

When samplePop is run on a set of simulated data of, say, 100 simulated trajectories (100 "subjects"), it will produce a matrix with 100 rows (100 "subjects"). But if it makes sense in the context of your problem (e.g., multiple samples per patient?) you can of course run samplePop repeatedly.

# 7.2 Differences between "samplePop" and "oncoSimulSample"

samplePop provides two sampling times: "last" and "uniform". It also allows you to sample at the first sample time(s) at which the population(s) reaches a given size, which can be either the same or different for each simulation (with argument popSizeSample). "last" means to sample each individual in the very last time period of the simulation. "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.

With oncoSimulSample we obtain samples that correspond to timeSample = "last" in samplePop by specifying a unique value for detectionSize and detectionDrivers. The data from each simulation will correspond to the time point at which those are reached (analogous to timeSample = "last"). How about uniform sampling? We pass a vector of detectionSize and detectionDrivers, where each value of the vector comes from a uniform distribution. This is not identical to the "uniform" sampling of oncoSimulSample, as we are not sampling uniformly over all time periods, but are stopping at uniformly distributed values over the stopping conditions. Arguably, however, the procedure in samplePop might be closer to what we mean with "uniformly sampled over the course of the disease" if that course is measured in terms of drivers or size of tumor.

An advantage of oncoSimulSample is that we can specify arbitrary sampling schemes, just by passing the appropriate vector detectionSize and detectionDrivers. A disadvantage is that if we change the stopping conditions we can not just resample the data, but we need to run it again.

There is no difference between oncoSimulSample and oncoSimulPop + samplePop in terms of the typeSample argument (whole tumor or single cell).

Finally, there are some additional differences between the two functions. oncoSimulPop can run parallelized (it uses mclapply). This is not done with oncoSimulSample because this function is designed for simulation experiments where you want to examine many different scenarios simultaneously. Thus, we provide additional stopping criteria (max.wall.time.total and max.num.tries.total) to determine whether to continue running the simulations, that bounds the total running time of all the simulations in a call to oncoSimulSample. And, if you are running multiple different scenarios, you might want to make multiple, separate, independent calls (e.g., from different R processes) to oncoSimulSample, instead of relying in mclapply, since this is likely to lead to better usage of multiple cores/CPUs if you are examining a large number of different scenarios.

## 8 Showing the genealogical relationships of clones

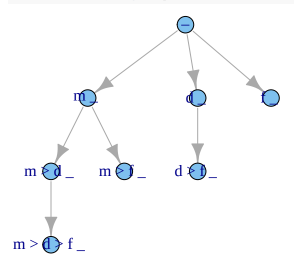
If you run simulations with keepPhylog = TRUE, the simulations keep track of when every clone is generated, and that will allow us to see the parent-child relationships between clones. (This is disabled by default).

Let us re-run a previous example:

```
set.seed(15)
tmp <- oncoSimulIndiv(examplesFitnessEffects[["o3"]],</pre>
                       model = "McFL",
                       mu = 5e-5,
                       detectionSize = 1e8,
                       detectionDrivers = 3,
                       sampleEvery = 0.025,
                       max.num.tries = 10,
                       keepEvery = 5,
                       initSize = 2000,
                       finalTime = 20000,
                       onlyCancer = FALSE,
                       extraTime = 1500,
                       keepPhylog = TRUE
tmp
##
## Individual OncoSimul trajectory with call:
    oncoSimulIndiv(fp = examplesFitnessEffects[["o3"]], model = "McFL",
       mu = 5e-05, detectionSize = 1e+08, detectionDrivers = 3,
##
       sampleEvery = 0.025, initSize = 2000, keepEvery = 5, extraTime = 1500,
##
##
       finalTime = 20000, onlyCancer = FALSE, keepPhylog = TRUE,
       max.num.tries = 10)
##
##
##
     NumClones TotalPopSize LargestClone MaxNumDrivers MaxDriversLast
                                                       3
## 1
             8
                        4071
                                     2632
                                                                        3
     {\it NumDriversLargestPop\ TotalPresentDrivers\ FinalTime\ NumIter}
##
                                                    6259 251954
                         3
## 1
                                              3
     HittedWallTime HittedMaxTries errorMF minDMratio minBMratio
##
## 1
              FALSE
                              FALSE 0.01442
                                                   6161
                                                               6667
##
     OccurringDrivers
## 1
              d, f, m
##
## Final population composition:
##
         Genotype
                      N
## 1
             d_{-}
## 2
                      0
## 3
         d > f
                      0
## 4
                      0
             f_{-}
                      0
## 5
```

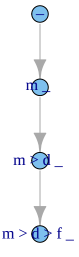
We can plot the parent-child relationships<sup>14</sup> of every clone ever created (with fitness larger than 0—clones without viability are never shown):

### plotClonePhylog(tmp, N = 0)



However, we often only want to show clones that exist (have number of cells > 0) at a certain time (while of course showing all of their ancestors, even if those are now extinct —i.e., regardless of their current numbers).

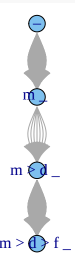
#### plotClonePhylog(tmp, N = 1)



If we set keepEvents = TRUE the arrows show how many times each clone appeared:
(The next can take a while)

<sup>&</sup>lt;sup>14</sup>There are several packages in R devoted to phylogenetic inference and related issues. For instance, *ape*. I have not used that infrastructure because of our very specific needs and circumstances; for instance, internal nodes are observed, we can have networks instead of trees, and we have no uncertainty about when events occurred.

### plotClonePhylog(tmp, N = 1, keepEvents = TRUE)



And we can show the plot so that the vertical axis is proportional to time (though you might see overlap of nodes if a child node appeared shortly after the parent):

```
plotClonePhylog(tmp, N = 1, timeEvents = TRUE)
```



We can obtain the adjacency matrix doing

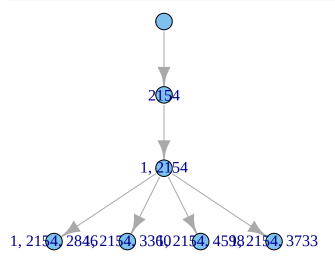
We can see another example here:

```
set.seed(456)
mcf1s <- oncoSimulIndiv(mcf1,</pre>
```

```
model = "McFL",
mu = 1e-7,
detectionSize = 1e8,
detectionDrivers = 100,
sampleEvery = 0.025,
keepEvery = 2,
initSize = 2000,
finalTime = 1000,
onlyCancer = FALSE,
keepPhylog = TRUE)
```

Showing only clones that exist at the end of the simulation (and all their parents):

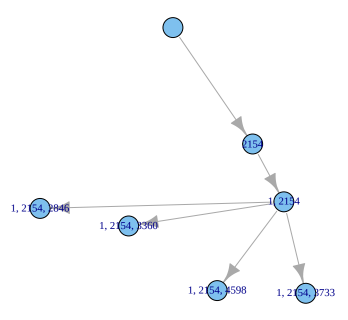
### plotClonePhylog(mcf1s, N = 1)



Notice that the labels here do not have a "\_\_", since there were no order effects in fitness. However, the labels show the genes that are mutated, just as before.

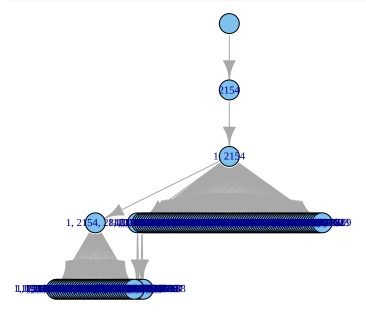
Similar, but with vertical axis proportional to time:

```
par(cex = 0.7)
plotClonePhylog(mcf1s, N = 1, timeEvents = TRUE)
```



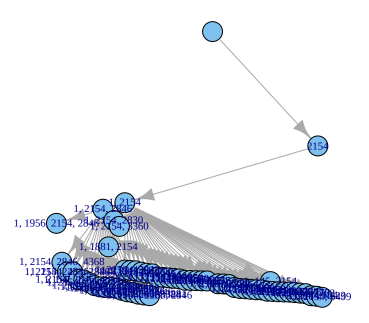
What about those that existed in the last 200 time units?

```
par(cex = 0.7)
plotClonePhylog(mcf1s, N = 1, t = c(800, 1000))
```



And try now to show also when the clones appeared (we restrict the time to between 900 and 1000, to avoid too much clutter):

```
par(cex = 0.7)
plotClonePhylog(mcf1s, N = 1, t = c(900, 1000), timeEvents = TRUE)
```

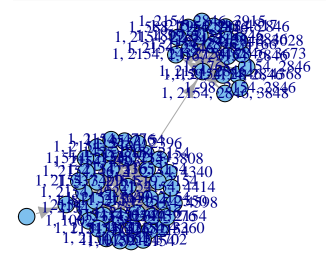


(By playing with t, it should be possible to obtain animations of the phylogeny. We will not pursue it here.)

If the previous graph seems cluttered, we can represent it in a different way by calling *igraph* directly after storing the graph and using the default layout:

```
g1 <- plotClonePhylog(mcf1s, N = 1, t = c(900, 1000),
returnGraph = TRUE)
```

### plot(g1)



which might be easier to show complex relationships or identify central or key clones.

It is of course quite possible that, especially if we consider few genes, the parent-child relationships will form a network, not a tree, as the same child node can have multiple parents. You can play with this example, modified from one we saw before (section 3.4.6):

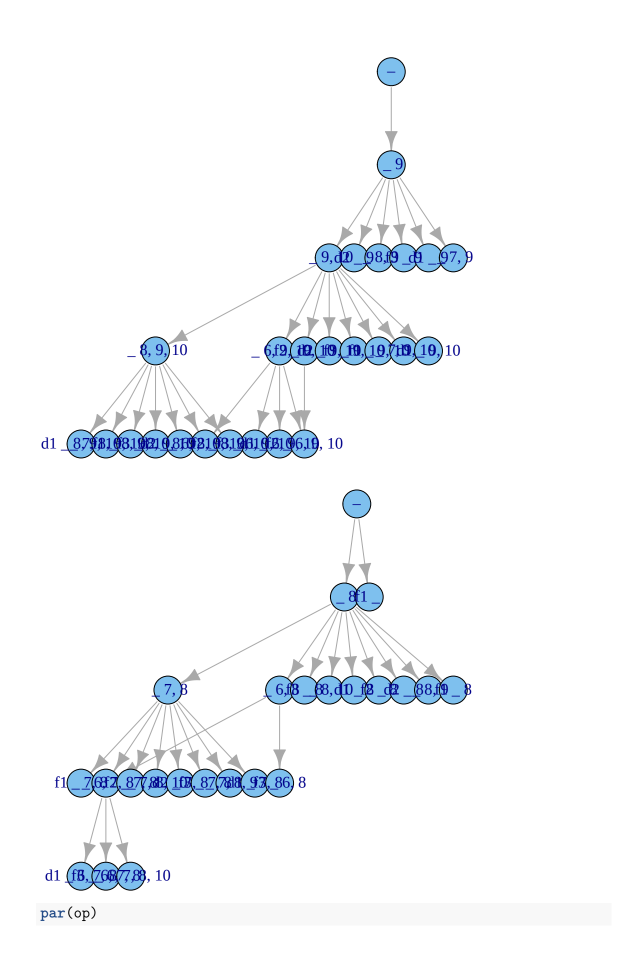
```
op <- par(ask = TRUE)
while(TRUE) {</pre>
```

### 8.1 Parent-child relationships from multiple runs

If you use oncoSimulPop you can store and plot the "phylogenies" of the different runs:

We will plot the first two:

```
op <- par(mar = rep(0, 4), mfrow = c(2, 1))
plotClonePhylog(oiP1[[1]])
plotClonePhylog(oiP1[[2]])</pre>
```



This is so far disabled in function <code>oncoSimulSample</code>, since that function is optimized for other uses. This might change in the future.

## 9 Generating random fitness landscapes

## 9.1 Random fitness landscapes from a Rough Mount Fuji model

In most of the examples seen above, we have fully specified the fitness of the different genotypes (either by providing directly the full mapping genotypes to fitness, or by providing that mapping by specifying the effects of the different gene combinations). In some cases, however, we might want to specify a particular model that generates the fitness landscape, and then have fitnesses be random variables obtained under this model. In other words, in this random fitness landscape the fitness of the genotypes is a random variable generated under some specific model. Random fitness landscapes are used extensively, for instance, to understand the evolutionary consequences of different types of epistatic interactions (e.g., Szendro, Schenk, et al. 2013; Franke et al. 2011) and there are especially developed tools for plotting and analyzing random fitness landscapes (e.g., Brouillet et al. 2015).

With OncoSimulR it is possible to generate mappings of genotype to fitness using the function rfitness that allows you to use from a pure House of Cards model to a purely additive model (see 9.2 for NK model). I have followed Szendro, Schenk, et al. (2013) and Franke et al. (2011) and model fitness as

$$f_i = -c \ d(i, reference) + x_i \tag{2}$$

where d(i,j) is the Hamming distance between genotypes i and j (the number of positions that differ), c is the decrease in fitness of a genotype per each unit increase in Hamming distance from the reference genotype, and  $x_i$  is a random variable (in this case, a normal deviate of mean 0 and standard deviation sd). You can change the reference genotype to any of the genotypes: for the deterministic part, you make the fittest genotype be the one with all positions mutated by setting reference = "max", or use the wildtype by using a string of 0s, or randomly select a genotype as a reference by using reference = "random" or reference = "random2".

And by changing c and sd you can flexibly modify the relative weight of the purely House of Cards vs. additive component. The expression used above is also very similar to the one on Greene and Crona (2014) if you use rfitness with the argument reference = "max".

What can you do with these genotype to fitness mappings? You could plot them, you could use them as input for oncoSimulIndiv and related functions, or you could export them (to\_Magellan) and plot them externally (e.g., in MAGELLAN: http://wwwabi.snv.jussieu.fr/public/Magellan/, Brouillet et al. (2015)).

```
## A small example

rfitness(3)

## A B C Fitness

## [1,] 0 0 0 1.000e+00

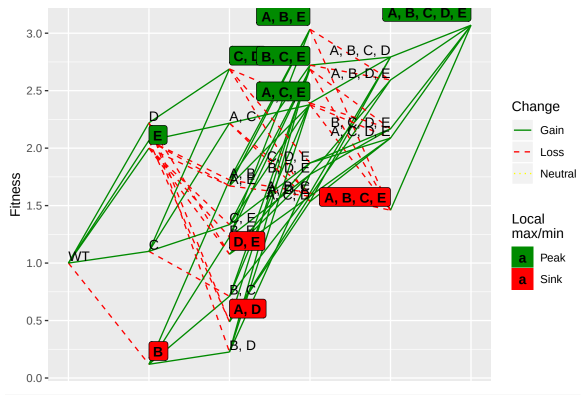
## [2,] 1 0 0 1.251e-10

## [3,] 0 1 0 4.814e-10
```

```
## [4,] 0 0 1 2.004e-01
## [5,] 1 1 0 1.733e-10
## [6,] 1 0 1 3.612e-02
## [7,] 0 1 1 1.214e-01
## [8,] 1 1 1 9.092e-02
## attr(,"class")
## [1] "matrix" "genotype_fitness_matrix"

## A 5-gene example, where the reference genotype is the
## one with all positions mutated, similar to Greene and Crona,
## 2014. We will plot the landscape and use it for simulations
## We downplay the random component with a sd = 0.5

r1 <- rfitness(5, reference = rep(1, 5), sd = 0.6)
plot(r1)</pre>
```



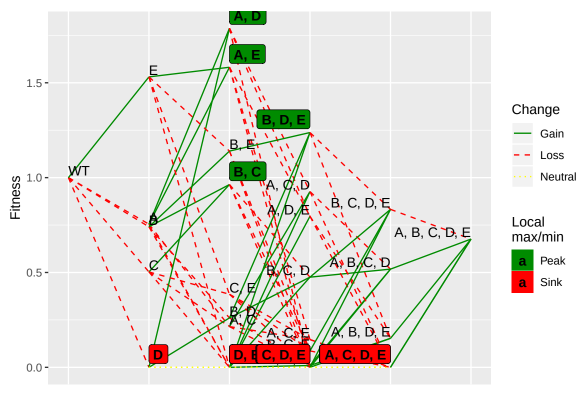
```
oncoSimulIndiv(allFitnessEffects(genotFitness = r1))
##
## Individual OncoSimul trajectory with call:
    oncoSimulIndiv(fp = allFitnessEffects(genotFitness = r1))
##
##
    NumClones TotalPopSize LargestClone MaxNumDrivers MaxDriversLast
                                   2e+08
                                                      0
## 1
                      2e+08
##
    NumDriversLargestPop TotalPresentDrivers FinalTime NumIter
## 1
                                            0
   HittedWallTime HittedMaxTries errorMF minDMratio minBMratio
```

```
## 1
               FALSE
                                FALSE
                                            NA
                                                    112583
                                                                 2e+05
##
     OccurringDrivers
## 1
##
## Final population composition:
     Genotype
                        N
## 1
                     1994
## 2
         A, D
                      240
             В
## 3
                        0
         B, D
                      210
## 4
         C, D
## 5
                    14170
             D 200020360
## 6
## 7
         D, E
                      438
```

### 9.2 Random fitness landscapes from Kauffman's NK model

You can also use Kauffman's NK model (e.g., Ferretti et al. 2016; Brouillet et al. 2015). We call the function fl\_generate from MAGELLAN (Brouillet et al. 2015).

```
rnk <- rfitness(5, K = 3, model = "NK")
plot(rnk)</pre>
```



```
oncoSimulIndiv(allFitnessEffects(genotFitness = rnk))
##
## Individual OncoSimul trajectory with call:
## oncoSimulIndiv(fp = allFitnessEffects(genotFitness = rnk))
```

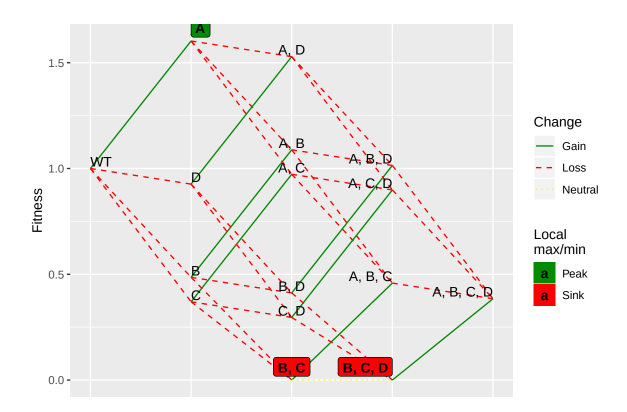
```
##
##
     NumClones TotalPopSize LargestClone MaxNumDrivers MaxDriversLast
## 1
                  107811448
                               107808707
     NumDriversLargestPop TotalPresentDrivers FinalTime NumIter
##
## 1
                                              0
                                                      852
     HittedWallTime HittedMaxTries errorMF minDMratio minBMratio
##
## 1
              FALSE
                             FALSE
                                         NA
                                                 163291
                                                             2e+05
     {\it OccurringDrivers}
##
## 1
##
## Final population composition:
     Genotype
                      N
## 1
                    463
## 2
                    1627
         A, E
## 3
            B
                      0
## 4
         B, E
                    494
## 5
            C
                      0
## 6
         C, E
                     157
            E 107808707
## 7
```

### 9.3 Random fitness landscapes from an additive model

This model evaluates fitness with different contributions of each allele, which will be randomly generated.

Given a number a genes by the user, the code uses rnorm to generate random contribution for the mutated allele in each locus. Later, this constributions will be used in the generation of the matrix that gives the value of fitness for each combination of wild type/mutated alleles by addition of the values for each locus and combination.

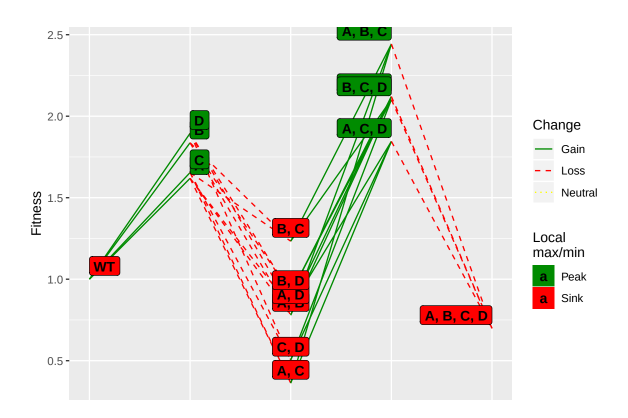
```
radd <- rfitness(4, model = "Additive", mu = 0, sd = 0.5)
plot(radd)</pre>
```



## 9.4 Random fitness landscapes from Eggbox model

You can also use Eggbox model (e.g., Ferretti et al. 2016; Brouillet et al. 2015), where each locus is either high or low fitness (depending on the "e" parameter value), with a systematic change between each neighbor. We call the function fl\_generate from MAGELLAN (Brouillet et al. 2015) to generate these landscapes.

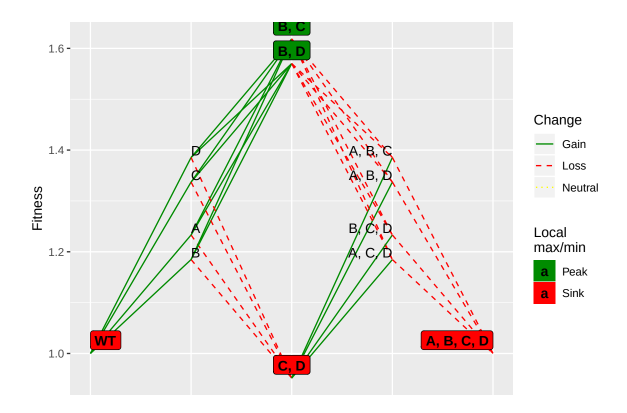
```
regg <- rfitness(4, model = "Eggbox", e = 1, E = 0.5)
plot(regg)</pre>
```



## 9.5 Random fitness landscapes from Ising model

In the Ising model (e.g., Ferretti et al. 2016; Brouillet et al. 2015), loci are arranged sequentially and each locus interacts with its physical neighbors. For each pair of interacting loci, there is a cost to (log)fitness if both alleles are not identical (and therefore 'compatible'); in this case, the cost for incompatibility i is applied. The last and the first loci will interact only if 'circular' is set. The implementation of this model is decribed (Brouillet et al. 2015), and we use a call to MAGELLAN code to generate the landscape.

```
ris <- rfitness(g = 4, model = "Ising", i = 0.002, I = 0.45)
plot(ris)
```

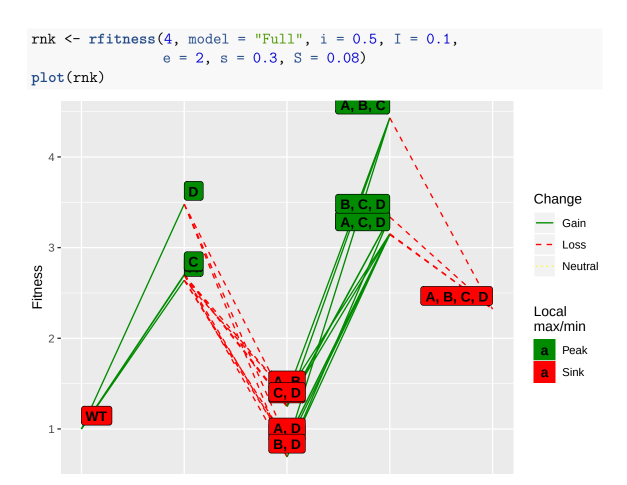


### 9.6 Random fitness landscapes from Full models

MAGELLAN also offers the possibility to combine different models with their own parameters in order to generate a Full model. The models combined are:

- House of cards: H is the number of interacting genes.
- Multiplicative (what we call additive): s and S mean and SD for generating random fitnesses. d is a diminishing (negative) or increasing (positive) return as you approach the peak.
- Kauffman NK: each locus interacts with any other K loci that can be chosen randomly pasing "r = TRUE" or among its neighbors.
- RMF: explained at 9.1.
- Ising: i and I mean and SD for incompatibility. If circular option is provided, the last and first alleles can interact (circular arrangement).
- Eggbox: In this model, each locus is considered as high or low fitness. From one locus to another the fitness changes its sign so that one is benefficial and its neighbor is detrimental. e and E are fitness and noise for fitness.
- Optimum: there is an optimum fitness contribution defined by o mu and O sigma and every locus has a production pand P (also mean and sd respectively).

All models can be taken into account for the fitness calculation. With default parameters, neither of Ising, Eggbox or Optimum contribute to fitness lanadcape generation as all i, e, o and p all == 0. Also, as all parameters referring to standard deviations have value == -1, those are also have no effect unless changed. Further details can be found in MAGELLAN's webpage http://wwwabi.snv.jussieu.fr/public/Magellan/ and (Brouillet et al. 2015).



## 9.7 Epistasis and fitness landscape statistics

We can call MAGELLAN's (Brouillet et al. 2015) fl\_statistics to obtain fitness landscape statistics, including measures of sign and reciprocal sign epistasis. See the help of Magellan\_stats for further details on output format. For example:

<pre>rnk1 &lt;- rfitness(6, K = 1, model = "NK")</pre>								
Magellan_stats(rnk1)								
##	ngeno	npeaks	nsinks	gamma	gamma .	r.s		
##	64.000	1.000	5.000	0.876	0.700	0.644		
##	nchains	nsteps	nori	depth	magn	sign		
##	1.000	9.000	6.000	4.000	0.438	0.150		
##	rsign	f.1 .	X.2.	f.3.	${\it mode}_{\_}f$	$outD\_m$		
##	0.075	0.843	0.149	0.007	1.000	2.107		
##	$\mathit{outD}\_\mathit{v}$	$steps\_{\it m}$	$reach\_{\it m}$	$fitG\_m$	$opt\_i$	${\it mProbOpt\_O}$		
##	2.095	3.329	12.048	32.000	3.000	1.000		
<pre>rnk2 &lt;- rfitness(6, K = 4, model = "NK")</pre>								
rnk2	<- rfitnes	s(6, K = 4,	model = "NK	")				
	<pre>&lt;- rfitnes: llan_stats(</pre>	•	model = "NK	")				
		•	model = "NK  nsinks	") gamma	gamma.	r.s		
Mage	llan_stats(	rnk2)			gamma. 0.138	r.s 3.289		

##	5.000	8.000	7.000	2.000	0.392	0.354
##	rsign	f.1.	X.2.	f.3	$mode_f$	•
##	0.254	0.182	0.282	0.536	3.000	2.212
##	$\mathit{outD}\_\mathit{v}$	$steps\_{\it m}$	$reach\_m$	$fitG\_m$	$opt\_i$	$mProbOpt_0$
##	3.460	2.028	13.684	34.842	14.000	0.205
##	$opt\_i.1$	${\it mProbOpt\_1}$	$opt\_i.2$	${\it mProbOpt\_2}$	$opt\_i.3$	${\it mProbOpt\_3}$
##	18.000	0.055	23.000	0.244	28.000	0.087
##	$opt_i.4$	$mProbOpt\_4$	$opt\_i.5$	${\it mProbOpt\_5}$	$opt\_i.6$	${\it mProbOpt\_6}$
##	32.000	0.104	35.000	0.179	52.000	0.126

(These fitness landscapes are, of course, frequency-independent fitness landscapes; with frequency-dependent fitness, as in section  $@ref\{fdf\}$  fitness landscapes as such are not defined.)

## 10 Frequency-dependent fitness

NOTE: This functionality is only available in the freq-dep-fitness branch See: https://github.com/rdiaz02/OncoSimul/tree/freq-dep-fitness. (Note also that except for the example of 10.2, based on Hurlbut et al. (2018), the examples below are not used because of their biological realism, but rather to show some key features of the software)

With frequency-dependence fitness we can make fitness (actually, birth rate) depend on the frequency of other genotypes. We specify how the fitness (birth rate) of each genotype depends (or not) on other genotypes. Thus, this is similar to the explicit mapping of genotypes to fitness (see 3.1.1), but fitness can be a function of the abundance (relative or absolute) of other genotypes. Frequency-dependent fitness allows you to examine models from **game theory** and **adaptive dynamics**. Game theory has long tradition in evolutionary biology (Maynard Smith 1982) and has been widely used in cancer (see, for example, Tomlinson 1997; Basanta and Deutsch 2008; Archetti and Pienta 2019, for classical papers that cover from early uses to a very recent review).

Since birth rate can be an arbitrary function of the frequencies of other clones, we can model competition, cooperation and mutualism, parasitism and predation, and commensalism. (Recall that in the "Exp" model death rate is constant and fixed to 1. In the "McFL" and "McFLD" models, death rate is density-dependent —but not frequency-dependent. You can thus model all those phenomena by, for example, making the effects of clones i, j on each other and on their own be asymmetric on their birth rates). See examples in section 10.4; as explained there, if you use the "Exp" model, you might want to decrease the value of sampleEvery.

The procedure for working with the frequency-dependent functionality is the general one with OncoSimulR. We first create a data frame with the mapping between genotypes and their (frequency-dependent) fitness, similar to section 3.1.1. For example, a two-column data frame, where the first column are the genotypes and the second column contains, as strings, the expressions for the function that relate fitness to frequencies of other genotypes. (We can also use a 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 expressions for the fitness specifications; see oncoSimulIndiv and allFitnessEffects for examples). Once this data frame is created, we pass it to allFitnessEffects. From there, simulation proceeds as usual.

How complex can the functions that specify fitness be? We use library ExprTk for the fitness specifications so the range of functions you can use is very large (http://www.partow.net/programming/exprtk/), including of course the usual arithmetic expressions, logical expressions (so you can model thresholds or jumps and use step functions), and a wide range of mathematical functions (so linear, non-linear, convex, concave, etc, functions can be used, including of course affine fitness functions as in Gerstung, Nakhoul, and Beerenwinkel 2011).

### 10.1 A first example with frequency-dependent fitness

The following is an arbitrary example. We will model birth rate of some genotypes as a function of the relative frequencies of other genotypes; we use f\_1 to denote the relative frequency of the genotype with the first gene mutated, f\_1\_2 to denote the relative frequency of the genotype with the first and second genes mutated, etc, and f\_ to denote the frequency of the WT genotype —below, in sections 10.3, 10.5, and 10.4, we will use absolute number of cells instead of relative frequencies).

As you can see below, the birth rate of genotype "A" =  $1.2 + 1.5 * f_1_2$  and that of the wildtype =  $1 + 1.5 * f_1_2$ . Genotype "A, B" in this example could be a genotype whose presence leads to an increase in the growth of other genotypes (maybe via diffusible factors, induction of angiogenesis, etc). Genotype "B" does not show frequency-dependence. The birth rate of genotype "C" increases with the frequency of  $f_1_2$  and increases (adding 0.7) with the frequency of genotypes "A" and "B", but only if the sum of the frequencies of genotypes "A" and "B" is larger than 0.3. For genotype "A, B" its fitness increases with the square root of the sum of the frequencies of genotypes "A", "B", and "C", but it decreases (i.e., shows increased intra-clone competition) if its own frequency is larger than 0.5. Genotypes not defined explicitly have a fitness of 0.

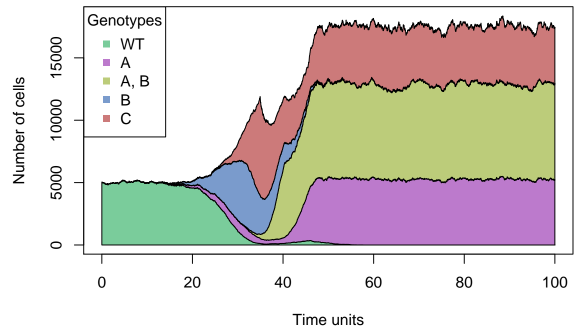
(In the data frame creation, we use stringsAsFactors = FALSE to avoid messages about conversions between factors and characters).

Let us verify that we have specified what we think we have specified using evalAllGenotypes (we have done this repeatedly in this vignette, for example in 1.3.2 or 1.6 or 5.1. Because of the way the code works, we need to pass the populations sizes at which we want fitness evaluated in evalAllGenotypes (this we do now in allFitnessEffects, but this might be moved to evalAllGenotypes in the future).

Note that when calling allFitnessEffects we have to set the parameter frequencyDependentFitness to TRUE. Since we are using relative frequencies, we also specify freqType = "rel" (the need to do this might be removed in the future). We will see below (10.3, 10.5, and 10.4) several examples with absolute numbers.

```
spPopSizes = c(100, 20, 20, 30, 0)))
##
     Genotype Fitness
## 1
          WT
                1.000
## 2
           \boldsymbol{A}
                1.300
## 3
            B
                1.400
            C 1.100
## 4
         A, B
## 5
                1.842
## 6
         A, C
                0.000
## 7
         B, C
                0.000
## 8 A, B, C
                0.000
evalAllGenotypes(allFitnessEffects(genotFitness = gffd,
                         frequencyDependentFitness = TRUE,
                         frequencyType = "rel",
                         spPopSizes = c(100, 30, 40, 0, 10)))
##
     Genotype Fitness
## 1
          WT
               1.083
## 2
                1.383
           \boldsymbol{A}
                1.400
## 3
            B
            C 1.856
## 4
        A, B
## 5
                1.824
         A, C
## 6
                0.000
## 7
         B, C
                0.000
## 8 A, B, C
                0.000
evalAllGenotypes(allFitnessEffects(genotFitness = gffd,
                         frequencyDependentFitness = TRUE,
                         frequencyType = "rel",
                         spPopSizes = c(100, 30, 40, 0, 100)))
##
     Genotype Fitness
## 1
          WT
               1.556
## 2
            A 1.856
## 3
            B
                1.400
## 4
            C 1.470
## 5
         A, B
                1.709
## 6
         A, C
                0.000
                0.000
## 7
         B, C
## 8 A, B, C
                0.000
```

The fitness specification is correct. Let us now create the allFitnessEffects object and simulate. We will use the McFL model, so in addition to the frequency dependence in the birth rates, there is also density dependence in the death rate (see section 3).



# 10.2 Hurlbut et al., 2018: a four-cell example with angiogenesis and cytotoxicity

The following example is based on Hurlbut et al. (2018). As explained in p. 3 of that paper, "Stromal cancer cells (A-) [WT in the code below] have no particular benefit or cost unique to themselves, and they are considered a baseline neutral cell within the context of the model. In contrast, angiogenesis-factor producing cells (A+) [A in the code below] vascularize the local tumor area which consequently introduces a nutrient rich blood to the benefit of all interacting cells. Nutrient recruitment expands when A+ cells interact with one another. Cytotoxic cells (C) release a chemical compound which harms heterospecific cells and increases their rate of cell death. The cytotoxic cells benefit from the resulting disruption in competition caused by the interaction. For simplicity, our model presumes that cytotoxic cells are themselves immune to

this class of agent. Finally, proliferative cells (P) possess a reproductive or metabolic advantage relative to the other cell types. In our model this advantage does not compound with the nutrient enrichment produced by vascularization when A+ cells are present; however, it does place the proliferative cell at a greater vulnerability to cytotoxins."

They provide, in p. 4, the payoff matrix reproduced in Figure 7:

Player\Opponent	<b>A</b> -	A+	P	С
A-	1	1+d	1	1-c
A+	1 - a + d	1 - a + d + f	1-a+d	1 - c - a + d
P	1 + g	1 + d + g	1 + g	(1+g)(1-c)
С	1 - b + e	1 - b + d + e	1 - b + e	1-b

Figure 7: Payoff matrix from Table 2 of Hurlbut et al., 2018, 'Game Theoretical Model of Cancer Dynamics withFour Cell Phenotypes', *Games*, 9, 61; doi:10.3390/g9030061.

As explained in p. 6 (equation 1) of Hurlbut et al. (2018), we can write the fitness of the four types as  $\mathbf{f} = G\mathbf{u}$ , where  $\mathbf{f}$  and  $\mathbf{u}$  are the vectors with the four fitnesses and frequencies (where each element of  $\mathbf{u}$  corresponds to the relative frequencies of stromal, angiogenic, proliferative, and cytotoxic cells), and G is the payoff matrix in Figure 7.

To allow modelling scenarios with different values for the parameters in Figure 7 we will define a function to create the data frame of frequency-dependent fitnesses. First, we will assume that each one of types A+, P, and C, are all derived from WT by a single mutation in one of three genes, say, A, P, C, respectively.

```
g, ") * f_3"),

paste0("1 - ", b, " + ",

e, " * f_ + ",

"(", d, " + ",

e, ") * f_1 + ",

e, " * f_2")),

stringsAsFactors = FALSE)
}
```

We can check we recover Figure 7:

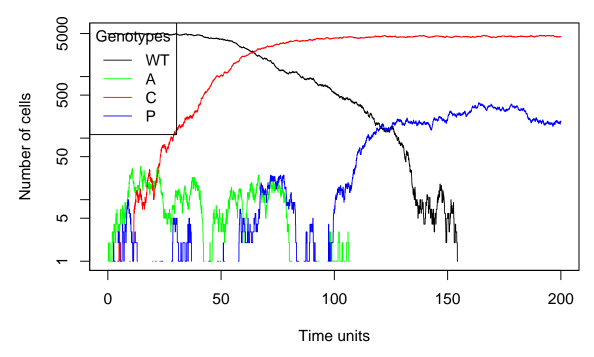
We could model a different set of ancestor-dependent relationships:

```
## Different assumption about origins from mutation:
## WT -> P; P -> A, P; P -> C, P
create_fe2 <- function(a, b, c, d, e, f, g,</pre>
                         gt = c("WT", "A", "P", "C", "A, P", "A, C",
                                  "C, P")) {
  data.frame(Genotype = gt,
             Fitness = c(
                  paste0("1 + ",
                        d, " * f_1_2 ",
                        "- ", c, " * f 2 3"),
                  "0",
                  paste0("1 + ", g, " + ",
                        d, " * f_1_2 ",
                        "- ", c, " * (1 + ",
                        g, ") * f_2_3"),
                  "0",
                  paste0("1 - ", a, " + ",
                  d, " + ",
                       f, " * f_1_2 ",
"- ", c, " * f_2_3"),
                  paste0("1 - ", b, " + ",
                        e, " * f_ + ",
                        "(", d, " + ",
                        e, ") * f_1_2 + ",
                        e, " * f 2")),
```

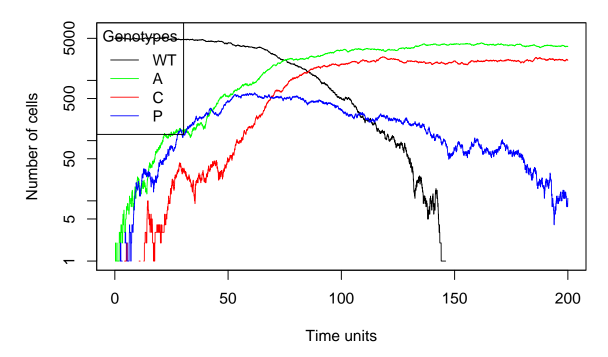
```
stringsAsFactors = FALSE)
}
## And check:
create_fe2("a", "b", "c", "d", "e", "f", "g")
## Genotype
                                                 Fitness
                              1 + d * f_1_2 - c * f_2_3
## 1
           WT
## 2
           \boldsymbol{A}
          P = 1 + g + d * f_1_2 - c * (1 + g) * f_2_3
## 3
## 4
          C
                  1 - a + d + f * f_1_2 - c * f_2_3
       A, P
## 5
        A, C
## 6
        C, P1 - b + e * f_ + (d + e) * f_1_2 + e * f_2
## 7
```

To show two examples, we will run the analyses Hurlbut et al. (2018) use for Figures 3a and 3b (p.8 of their paper):

```
## Figure 3a
afe_3_a <- allFitnessEffects(</pre>
                genotFitness =
                        create_fe(0.02, 0.04, 0.08, 0.06,
                                 0.15, 0.1, 0.06),
                frequencyDependentFitness = TRUE,
                frequencyType = "rel")
set.seed(2)
s_3_a <- oncoSimulIndiv(afe_3_a,</pre>
                     model = "McFL",
                      onlyCancer = FALSE,
                     finalTime = 200,
                     mu = 1e-4,
                     initSize = 5000,
                     keepPhylog = TRUE,
                      seed = NULL,
                      errorHitMaxTries = FALSE,
                     errorHitWallTime = FALSE)
## plot(s_3_a, show = "genotypes",
##
        xlim = c(40, 200),
        col = c("black", "green", "red", "blue"))
plot(s 3 a, show = "genotypes", type = "line",
     col = c("black", "green", "red", "blue"))
```

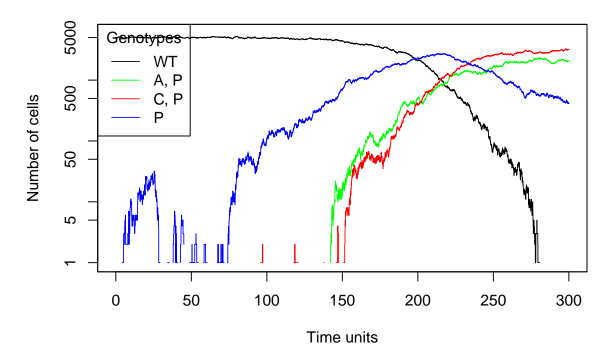


```
## Figure 3b
afe_3_b <- allFitnessEffects(</pre>
                genotFitness =
                        create_fe(0.02, 0.04, 0.08, 0.1,
                                  0.15, 0.1, 0.05),
                frequencyDependentFitness = TRUE,
                frequencyType = "rel")
set.seed(2)
s_3_b <- oncoSimulIndiv(afe_3_b,</pre>
                     model = "McFL",
                      onlyCancer = FALSE,
                     finalTime = 200,
                     mu = 1e-4,
                      initSize = 5000,
                     keepPhylog = TRUE,
                      seed = NULL,
                     errorHitMaxTries = FALSE,
                     errorHitWallTime = FALSE)
## plot(s_3_b, show = "genotypes",
        col = c("black", "green", "red", "blue"))
plot(s_3_b, show = "genotypes", type = "line",
     col = c("black", "green", "red", "blue"))
```



Of course, if we assume that the mutations leading to the different cell types are different, the results can change:

```
## Figure 3b. Now with WT -> P; P -> A,P; P -> C,P
afe_3_b_2 <- allFitnessEffects(</pre>
                   genotFitness =
                       create_fe2(0.02, 0.04, 0.08, 0.1,
                                  0.15, 0.1, 0.05),
                   frequencyDependentFitness = TRUE,
                   frequencyType = "rel")
set.seed(2)
s_3_b_2 <- oncoSimulIndiv(afe_3_b_2,</pre>
                    model = "McFL",
                    onlyCancer = FALSE,
                    finalTime = 300,
                    mu = 1e-4,
                    initSize = 5000,
                    keepPhylog = TRUE,
                    seed = NULL,
                    errorHitMaxTries = FALSE,
                    errorHitWallTime = FALSE)
plot(s_3_b_2, show = "genotypes", type = "line",
     col = c("black", "green", "red", "blue"))
```



(Examples for the remaining Figures 2 and 3 are provided, using also oncoSimulPop, in file 'inst/miscell/hurlbut-ex.R')

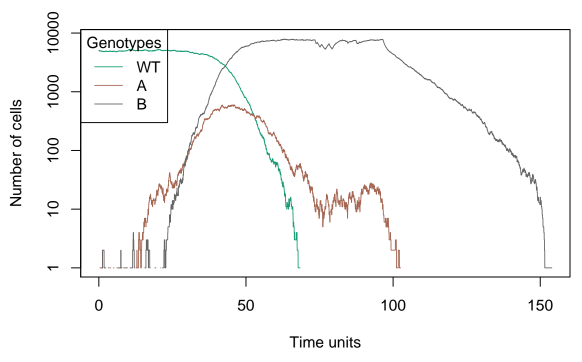
# 10.3 An example with absolute numbers and population collapse

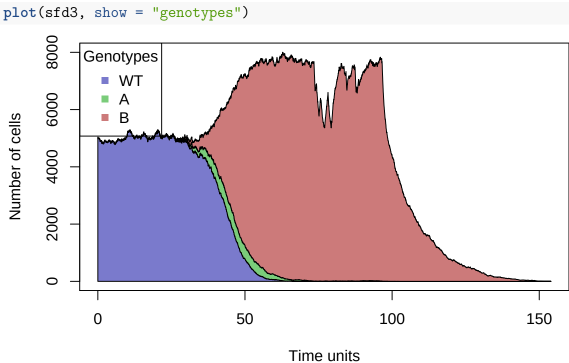
In the following example we use absolute numbers (thus the n\_1, etc, instead of the f\_1 in the fitness definition). This is a toy model where there is a change in fitness in two of the genotypes if the other is above a specified threshold (this also shows again the usage of a logical inequality). The genotype with gene B mutated has birth rate less than 1, unless there are at least more than 10 cells with genotype A mutated.

As usual, let us verify that we have specified what we think we have specified using evalAllGenotypes:

```
## 1
             WT
                     1.0
## 2
             \boldsymbol{A}
                     1.2
## 3
             B
                     0.9
                     0.0
## 4
          A, B
evalAllGenotypes(allFitnessEffects(genotFitness = gffd3,
                            frequencyDependentFitness = TRUE,
                            frequencyType = "abs",
                            spPopSizes = c(100, 11, 1)))
##
     Genotype Fitness
## 1
            WT
                     1.0
## 2
             \boldsymbol{A}
                     1.0
## 3
             B
                     1.3
                     0.0
          A, B
```

In this simulation, the population collapses: genotype B is able to invade the population when there are some A's around. But as soon as A disappears due to competition from B, B collapses as its birth rate becomes 0.9, less than the death rate; we are using the "McFLD" model (where death rate  $D(N) = \max(1, \log(1 + N/K))$ ); see details in 3.2.1.1.

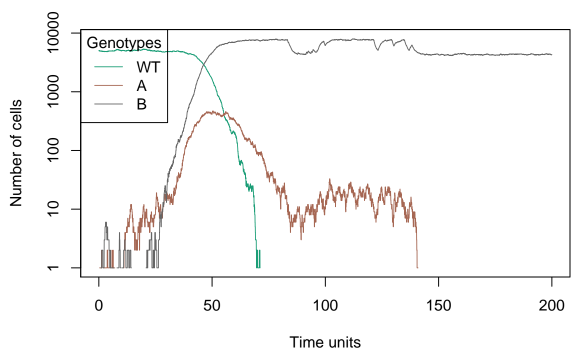


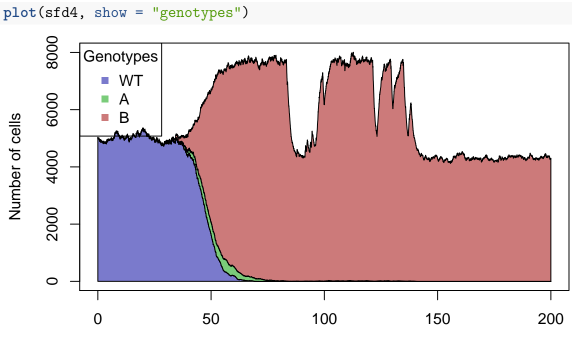


```
sfd3
##
## Individual OncoSimul trajectory with call:
## oncoSimulIndiv(fp = afd3, model = "McFLD", mu = 1e-04, initSize = 5000,
## finalTime = 200, onlyCancer = FALSE, keepPhylog = FALSE,
## errorHitWallTime = FALSE, errorHitMaxTries = FALSE, seed = NULL)
##
## NumClones TotalPopSize LargestClone MaxNumDrivers MaxDriversLast
```

```
## 1
     NumDriversLargestPop TotalPresentDrivers FinalTime NumIter
##
## 1
                         0
                                              0
                                                      154
    HittedWallTime HittedMaxTries errorMF minDMratio minBMratio
##
## 1
              FALSE
                              FALSE 0.00894
                                                   5000
                                                               5000
##
     OccurringDrivers
## 1
##
## Final population composition:
##
     Genotype N
## 1
              0
## 2
            A O
## 3
            B 0
```

Had we used the "usual" death rate expression, that can lead to death rates below 1 (see 3.2.1.1), we would have obtained a population that stabilizes around a final value slightly below the initial one (the one that corresponds to a death rate equal to 0.9):





```
sfd4
##
## Individual OncoSimul trajectory with call:
## oncoSimulIndiv(fp = afd3, model = "McFL", mu = 1e-04, initSize = 5000,
## finalTime = 200, onlyCancer = FALSE, keepPhylog = FALSE,
## errorHitWallTime = FALSE, errorHitMaxTries = FALSE, seed = NULL)
##
## NumClones TotalPopSize LargestClone MaxNumDrivers MaxDriversLast
```

Time units

```
0
## 1
                        4290
                                      4290
##
     NumDriversLargestPop TotalPresentDrivers FinalTime NumIter
## 1
                          0
                                               0
                                                       200
                                                               8154
##
     HittedWallTime HittedMaxTries errorMF minDMratio minBMratio
                                                                5000
   1
                               FALSE 0.01178
##
               FALSE
                                                    4843
     OccurringDrivers
##
## 1
##
## Final population composition:
##
     Genotype
                  N
## 1
                  0
## 2
                  0
             Α
            B 4290
## 3
## Check final pop size corresponds to birth = death
K < -5000/(exp(1) - 1)
K
## [1] 2910
log1p(4290/K)
## [1] 0.906
```

# 10.4 Predator-prey, commensalism, and consumer-resource models

Since birth rates can be arbitrary functions of frequencies of other clones, we can easily model classical ecological models, such as predator-prey, competition, commensalism and, more generally, consumer-resource models (see, for example, section 3.4 in Otto and Day 2007).

OncoSimulR was originally designed as a forward-time genetic simulator; thus, we need to use a simple trick to get the system going. For example, suppose we want to model a predator-prey system; we can do this having a WT that can mutate into either preys or predators. (This will not be necessary once we can specify starting the simulation from populations with arbitrary numbers of different clones.) There is, in fact, a small leakage in the system because both preys and predators are "leaking" a small number of children, via mutation to a non-viable predator-and-prey genotype, but this is negligible relative to their birth/death rates.

We will use below the usual consumer-resource model with Lotka-Volterra equations. To model them, we use the "Exp" model, which has a constant death rate of 1, and directly translate the usual expressions (e.g., expressions 3.15a and 3.15b in p. 73 of Otto and Day 2007) for rates of change into the birth rate. Of course, you can use any other function for how rates of change of each "species" depend on the numbers of the different species, including constant inflow and outflow, Type I, Type II, and Type III functional responses, etc. And you can model systems that involve multiple different types of preys, predators, commensals, etc (see the example in 10.2 for a four-type example).

What about sampleEvery? In "for real" work, and specially with complex models, you might want to decrease it, or at least examine how much results are affected by changes in sampleEvery. Decreasing sampleEvery will result in birth rates being updated more frequently, and we use the BNB algorithm, which updates all rates only when the whole population is sampled. (Recall that by default, in the McFL and McFLD models the setting for sampleEvery is smaller than in the Exp model). Further discussion of these issues is provided in sections 10.7 and 14.6.

Since the WT genotype is just a trick used to get the system going, we will make sure it disappears from the system soon after we get it to have both preys and predators (and we use the max function to prevent birth rate from ever becoming negative or identically 0). Finally, note that we can make this much more sophisticated; for example, we could get the system going differently by having different mutations from WT to prey and predator.

### 10.4.1 Competition

First, create a function that will generate the usual Lotka-Volterra expressions for competition models:

```
G fe LV <- function(r1, r2, K1, K2, a_12, a_21, awt = 1e-4,
                                 gt = c("WT", "S1", "S2")) {
    data.frame(Genotype = gt,
               Fitness = c(
                  paste0("max(0.1, 1 - ", awt, " * (n 2 + n 1))"),
                  paste0("1 + ", r1,
                         " * (1 - (n 1 + ", a 12, " * n 2)/", K1,
                  paste0("1 + ", r2,
                         " * ( 1 - (n_2 + ", a_21, " * n 1)/", K2,
                  ))
}
## Show expressions for birth rates
G_fe_LV("r1", "r2", "K1", "K2", "a_12", "a_21", "awt")
##
     Genotype
                   max(0.1, 1 - awt * (n 2 + n 1))
## 1
           S1 \ 1 + r1 * (1 - (n_1 + a_1 + a_2)/K1)
## 2
           S2\ 1 + r2 * (1 - (n_2 + a_21 * n_1)/K2)
## 3
```

Remember the above are the birth rates for a model with death rate = 1 (the "Exp" model). That is why we added a 1 to the birth rate. If you subtract 1 from the expressions for the birth rates of predators and prey above you get the standard expressions for the differential equations for Lotka-Volterra model of competition:  $\frac{\mathrm{d}n_1}{\mathrm{d}t} = r_1(1 - \frac{n_1 + \alpha_{12}n_2}{K_1}).$ 

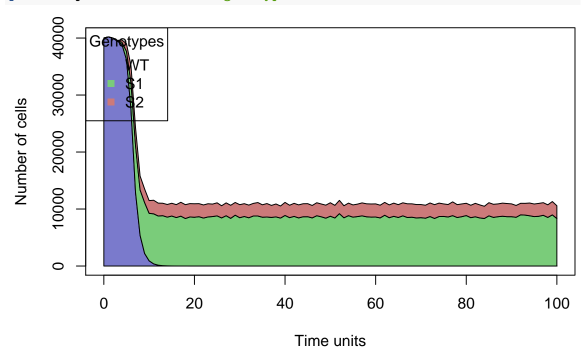
Now, run that model by setting appropriate parameters; see how we have a 12 > 0

```
and a_{21} > 0.
```

```
fe_competition <-</pre>
    allFitnessEffects(
        genotFitness =
            G_fe_LV(1.5, 1.4, 10000, 4000, 0.6, 0.2,
                   gt = c("WT", "S1", "S2")),
        frequencyDependentFitness = TRUE,
        frequencyType = "abs")
competition <- oncoSimulIndiv(fe_competition,</pre>
                             model = "Exp",
                             onlyCancer = FALSE,
                             finalTime = 100,
                             mu = 1e-4,
                              initSize = 40000,
                             keepPhylog = TRUE,
                             seed = NULL,
                              errorHitMaxTries = FALSE,
                              errorHitWallTime = FALSE)
```

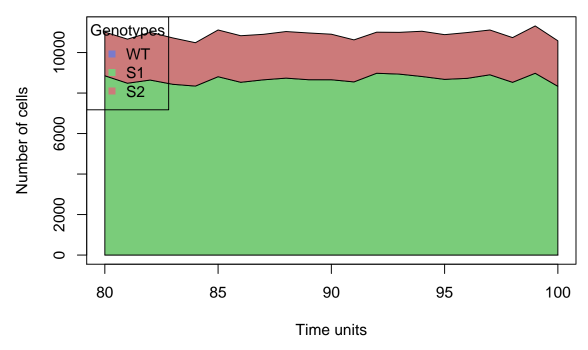
If we plot the whole simulation, we of course see the WT:

```
plot(competition, show = "genotypes")
```

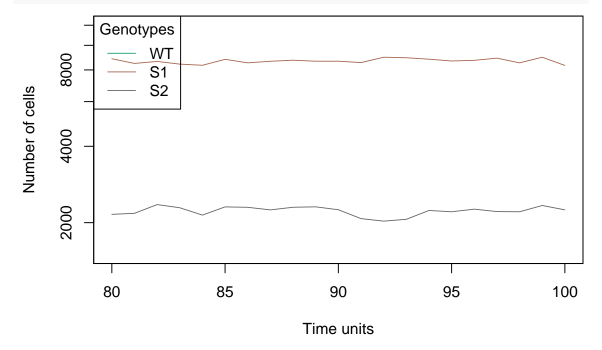


but we can avoid that by showing the plot after the WT are long gone:

```
plot(competition, show = "genotypes",
    xlim = c(80, 100))
```







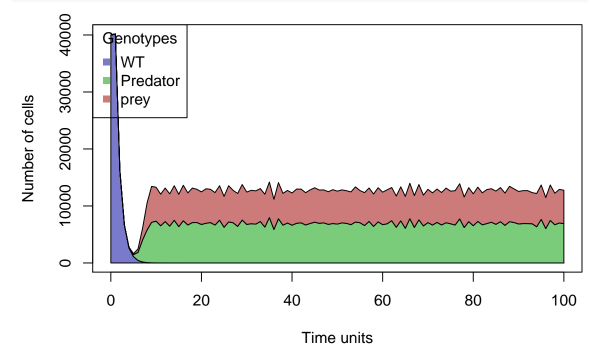
## 10.4.2 Predator-prey, first example

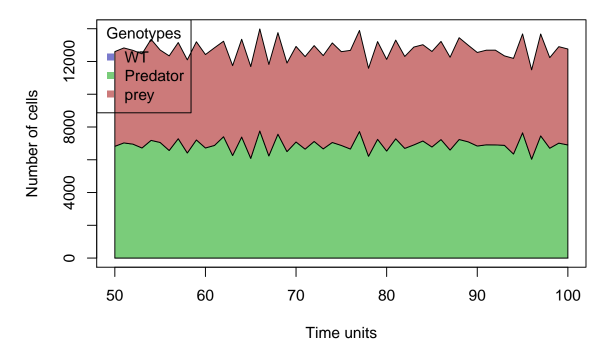
The simplest model would use the above Lotka-Volterra expressions and set one of the a\_ to be negative (see, for example, Otto and Day 2007, 73). Let's turn former "S2" and thus a\_21 < 0 (to make the effects more salient, we also increase that value in magnitude).

fe\_pred\_prey <-</pre>

```
allFitnessEffects(
        genotFitness =
            G_fe_LV(1.5, 1.4, 10000, 4000, 0.6, -0.5, awt = 1,
                  gt = c("WT", "prey", "Predator")),
        frequencyDependentFitness = TRUE,
        frequencyType = "abs")
## All single-gene genotypes as input to to_genotFitness_std
pred_prey <- oncoSimulIndiv(fe_pred_prey,</pre>
                             model = "Exp",
                             onlyCancer = FALSE,
                             finalTime = 100,
                             mu = 1e-4,
                             initSize = 40000,
                             keepPhylog = TRUE,
                             seed = NULL,
                             errorHitMaxTries = FALSE,
                             errorHitWallTime = FALSE)
```

plot(pred\_prey, show = "genotypes")

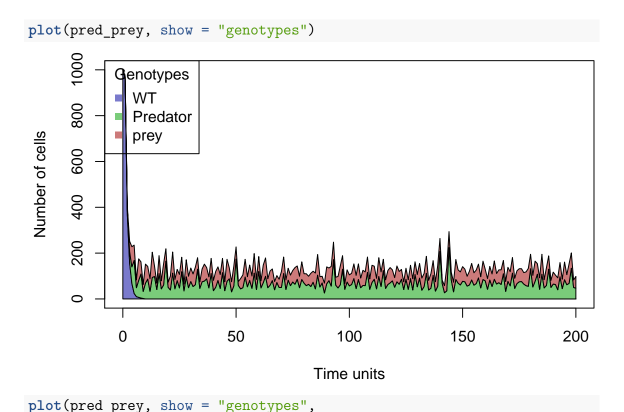


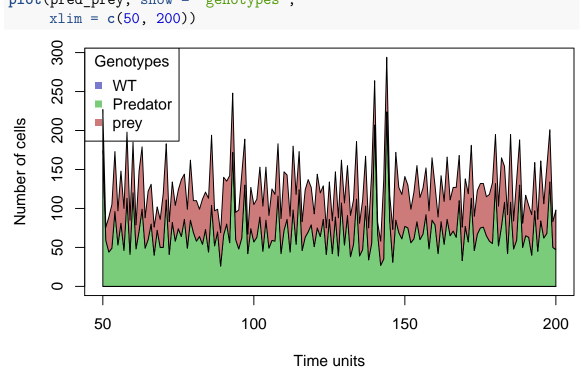


You can easily play with a range of parameters, say the carrying capacity of one of the species, to see how they affect the stochasticity of the system. Here we also start from a large number of WT (to generate preys and predators quickly) but we can set carrying capacity to much smaller numbers:

```
fe_pred_prey <-</pre>
    allFitnessEffects(
        genotFitness =
             G_fe_LV(1.5, 1.4, 100, 40, 0.6, -0.5, awt = 1,
                   gt = c("WT", "prey", "Predator")),
        frequencyDependentFitness = TRUE,
        frequencyType = "abs")
pred_prey <- oncoSimulIndiv(fe_pred_prey,</pre>
                              model = "Exp",
                              onlyCancer = FALSE,
                              finalTime = 200,
                             mu = 1e-3,
                              initSize = 1000,
                             keepPhylog = TRUE,
                              seed = NULL,
                              errorHitMaxTries = FALSE,
                              errorHitWallTime = FALSE)
```

Plotting the complete range shows the drop from the initial WT population size to the final carrying capacities (and the second plot zooms in after there are no WTs):





If you run the above model repeatedly, you will frequently find that only one of the species is left; and, yes, that could be the "Predator", as in the Lotka-Volterra expressions above there can be predators without prey. For example, you can check that the birth rate of the predator is larger than 1 even if there are 0 prey (identically 1 when  $n_2 = K_2$ :

```
evalAllGenotypes(allFitnessEffects(
   genotFitness =
       G_fe_LV(1.5, 1.4, 100, 40,
             0.6, -0.5, awt = 0.1,
             gt = c("WT", "prey", "Predator")),
   frequencyDependentFitness = TRUE,
   frequencyType = "abs",
   spPopSizes = c(0, 0, 20))
## All single-gene genotypes as input to to_genotFitness_std
          Genotype Fitness
##
## 1
                 WT
                      0.10
                      1.70
## 2
          Predator
## 3
              prey 2.32
## 4 Predator, prey 0.00
evalAllGenotypes(allFitnessEffects(
   genotFitness =
       G_fe_LV(1.5, 1.4, 100, 40,
             0.6, -0.5, awt = 0.1,
             gt = c("WT", "prey", "Predator")),
   frequencyDependentFitness = TRUE,
   frequencyType = "abs",
   spPopSizes = c(0, 0, 40)))
## All single-gene genotypes as input to to_genotFitness_std
          Genotype Fitness
## 1
                 WT
                     0.10
## 2
         Predator 1.00
## 3
                     2.14
              prey
## 4 Predator, prey
                      0.00
```

(Of course, with those population sizes, even if prey had a birth rate > 1, they cannot appear as there are no WT to generate prey).

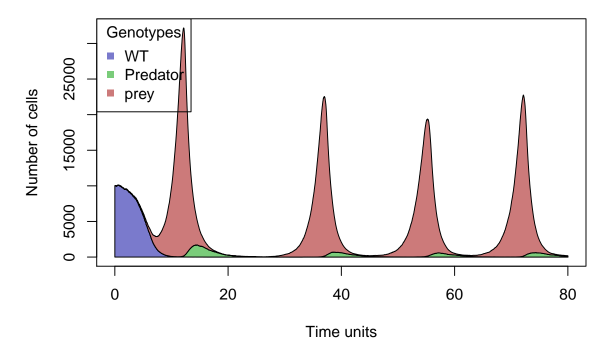
## 10.4.3 Predator-prey, second example

We use now the model in p. 76 of Otto and Day (2007), where prey grow exponentially in the absence of predators (and predators will eventually go extinct in the absence of prey):

$$\frac{\mathrm{d}n_1}{\mathrm{d}t} = r \ n_1 - a \ c \ n_1 \ n_2$$

$$\frac{\mathrm{d}n_2}{\mathrm{d}t} = \epsilon \ a \ c \ n_1 \ n_2 - \delta \ n_2$$

```
## Use e for epsilon and d for delta
C_fe_pred_prey <- function(r, a, c, e, d, awt = 0.1,</pre>
                            gt = c("WT", "prey", "Predator")) {
    data.frame(Genotype = gt,
               Fitness = c(
                   paste0("max(0.1, 1 - ", awt,
                          " * (n 2 + n 1))"),
                   paste0("1 + ", r, " - ", a,
                          " * ", c, " * n 2"),
                   paste0("1 + ", e, " * ", a,
                           " * ", c, " * n_1 - ", d)
               ))
}
C_fe_pred_prey("r", "a", "c", "e", "d")
     Genotype
                                        Fitness
## 1
           WT \max(0.1, 1 - 0.1 * (n_2 + n_1))
## 2
         prey
                           1 + r - a * c * n_2
## 3 Predator
                       1 + e * a * c * n_1 - d
fe_pred_prey2 <-</pre>
    allFitnessEffects(
        genotFitness =
            C_fe_pred_prey(r = .7, a = 1, c = 0.005,
                            e = 0.02, d = 0.4, awt = 0.001,
                            gt = c("WT", "prey", "Predator")),
        frequencyDependentFitness = TRUE,
        frequencyType = "abs")
## All single-gene genotypes as input to to_genotFitness_std
pred prey2 <- oncoSimulIndiv(fe pred prey2,</pre>
                             model = "Exp",
                             sampleEvery = 0.01,
                             mu = 1e-3,
                             onlyCancer = FALSE,
                             finalTime = 80,
                             initSize = 1e4,
                             keepPhylog = TRUE,
                             seed = NULL,
                             errorHitMaxTries = FALSE,
                             errorHitWallTime = FALSE)
plot(pred_prey2, show = "genotypes")
```



If you run that model repeatedly, sometimes the system will go extinct quickly, or you will only get prey growing exponentially.

## 10.4.4 Commensalism

Modelling commensalism simply requires changing the values of the  $\alpha$ , the a\_12 and a\_21. For example (not run, as this is just repetitive):

```
fe commens <-
    allFitnessEffects(
        genotFitness =
            G_fe_LV(1.2, 1.3, 5000, 20000,
                                  0, -0.2,
                                  gt = c("WT", "A", "Commensal")),
        frequencyDependentFitness = TRUE,
        frequencyType = "abs")
commens <- oncoSimulIndiv(fe_commens,</pre>
                             model = "Exp",
                             onlyCancer = FALSE,
                             finalTime = 100,
                             mu = 1e-4,
                             initSize = 40000,
                             keepPhylog = TRUE,
                             seed = NULL,
                             errorHitMaxTries = FALSE,
                             errorHitWallTime = FALSE)
plot(commens, show = "genotypes")
```

# 10.5 Frequency-dependent fitness: can I mix relative and absolute frequencies?

Yes, of course, since you can always use an absolute specification with the appropriate quotient. For example, the following two specifications are identical:

```
rar <- data.frame(Genotype = c("WT", "A", "B", "C"),</pre>
                 Fitness = c("1",
                             "1.1 + .3*f 2",
                             "1.2 + .4*f 1",
                             "1.0 + .5 * (f_1 + f_2)")
afear <- allFitnessEffects(genotFitness = rar,</pre>
                         frequencyDependentFitness = TRUE,
                         frequencyType = "rel",
                         spPopSizes = c(100, 200, 300, 400))
evalAllGenotypes(afear)
     Genotype Fitness
## 1
          WT
                 1.00
## 2
          \boldsymbol{A}
                 1.19
## 3
          B
                1.28
          C
## 4
                1.25
      A, B
## 5
              0.00
## 6
        A . C
                0.00
## 7
        B, C
               0.00
## 8 A, B, C
                 0.00
rar2 <- data.frame(Genotype = c("WT", "A", "B", "C"),</pre>
                 Fitness = c("1",
                             "1.1 + .3*(n 2/N)",
                             "1.2 + .4*(n_1/N)",
                             "1.0 + .5 * ((n 1 + n 2)/N)"))
afear2 <- allFitnessEffects(genotFitness = rar2,</pre>
                         frequencyDependentFitness = TRUE,
                         frequencyType = "abs",
                         spPopSizes = c(100, 200, 300, 400))
evalAllGenotypes(afear2)
##
     Genotype Fitness
## 1
           WT
                 1.00
## 2
         A 1.19
```

```
## 3 B
               1.28
## 4
           C
                1.25
## 5
       A, B
               0.00
## 6
        A, C
                0.00
## 7
        B, C
                0.00
## 8 A, B, C
                0.00
```

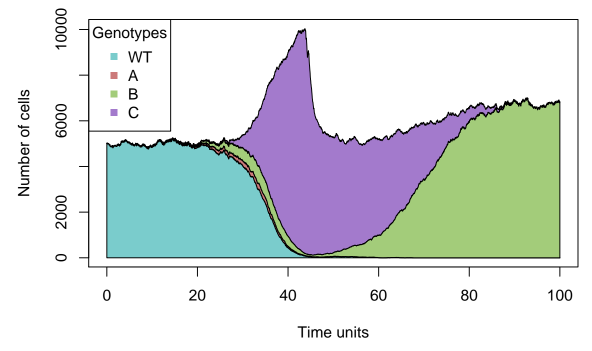
and simulating with them leads to identical results

```
set.seed(1)
tmp1 <- oncoSimulIndiv(afear,</pre>
                       model = "McFL",
                       onlyCancer = FALSE,
                       finalTime = 30,
                       mu = 1e-4,
                       initSize = 5000.
                       keepPhylog = FALSE,
                       seed = NULL,
                       errorHitMaxTries = FALSE,
                       errorHitWallTime = FALSE)
set.seed(1)
tmp2 <- oncoSimulIndiv(afear2,</pre>
                       model = "McFL",
                       onlyCancer = FALSE,
                       finalTime = 30,
                       mu = 1e-4,
                       initSize = 5000,
                       keepPhylog = FALSE,
                       seed = NULL,
                       errorHitMaxTries = FALSE,
                       errorHitWallTime = FALSE)
stopifnot(identical(print(tmp1), print(tmp2)))
##
## Individual OncoSimul trajectory with call:
    oncoSimulIndiv(fp = afear, model = "McFL", mu = 1e-04, initSize = 5000,
##
       finalTime = 30, onlyCancer = FALSE, keepPhylog = FALSE, errorHitWallTime =
##
       errorHitMaxTries = FALSE, seed = NULL)
##
##
    NumClones TotalPopSize LargestClone MaxNumDrivers MaxDriversLast
## 1
                       4988
                                   4661
##
    NumDriversLargestPop TotalPresentDrivers FinalTime NumIter
## 1
                        0
                                             0
    HittedWallTime HittedMaxTries errorMF minDMratio minBMratio
##
## 1
              FALSE
                        FALSE 0.006913
                                                   3259
                                                               3333
## OccurringDrivers
```

```
## 1
##
## Final population composition:
     Genotype
                 N
              4661
## 1
## 2
            A 319
## 3
            B
                 6
            C
                 2
## 4
##
## Individual OncoSimul trajectory with call:
    oncoSimulIndiv(fp = afear2, model = "McFL", mu = 1e-04, initSize = 5000,
       finalTime = 30, onlyCancer = FALSE, keepPhylog = FALSE, errorHitWallTime =
##
       errorHitMaxTries = FALSE, seed = NULL)
##
##
##
    NumClones TotalPopSize LargestClone MaxNumDrivers MaxDriversLast
             4
                       4988
                                    4661
##
    NumDriversLargestPop TotalPresentDrivers FinalTime NumIter
                                           0
## 1
                        0
                                                      30
                                                            1244
    HittedWallTime HittedMaxTries errorMF minDMratio minBMratio
##
## 1
                             FALSE 0.006913
                                                   3259
              FALSE
     OccurringDrivers
## 1
##
## Final population composition:
     Genotype
                N
## 1
              4661
## 2
            A 319
## 3
            В
                 6
## 4
            C
                 2
```

So you can always mix relative and absolute; here fitness of two genotypes depends on the relative frequencies of others, whereas fitness of the third on the absolute frequencies (number of cells):

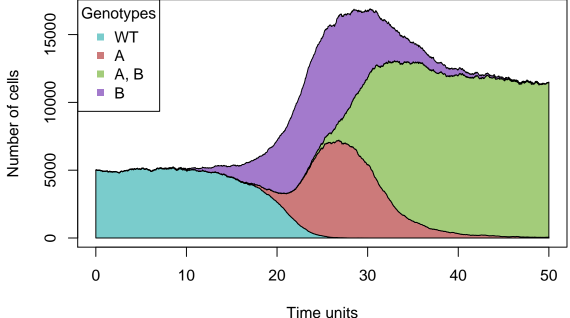
```
## 2
             \boldsymbol{A}
                   1.19
                   1.28
## 3
             B
             C
                   1.50
## 4
## 5
          A, B
                   0.00
          A, C
                   0.00
## 6
## 7
          B, C
                   0.00
## 8 A, B, C
                   0.00
set.seed(1)
tmp3 <- oncoSimulIndiv(afear3,</pre>
                         model = "McFL",
                         onlyCancer = FALSE,
                         finalTime = 100,
                         mu = 1e-4,
                         initSize = 5000,
                         keepPhylog = FALSE,
                         seed = NULL,
                         errorHitMaxTries = FALSE,
                         errorHitWallTime = FALSE)
plot(tmp3, show = "genotypes")
```

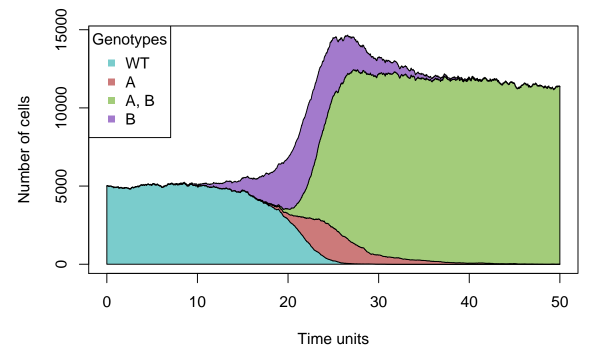


# 10.6 Frequency-dependent fitness: can I use genes with mutator effects?

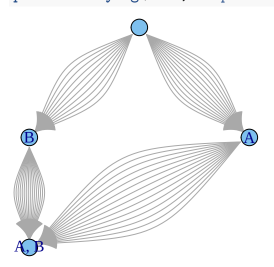
Yes. The following examples show it:

```
## Relative
r1fd <- data.frame(Genotype = c("WT", "A", "B", "A, B"),</pre>
                 Fitness = c("1",
                              "1.4 + 1*(f_2)",
                              "1.4 + 1*(f 1)",
                              "1.6 + f 1 + f 2"))
afe4 <- allFitnessEffects(genotFitness = r1fd,
                          frequencyDependentFitness = TRUE,
                          frequencyType = "rel")
set.seed(1)
s1fd <- oncoSimulIndiv(afe4,</pre>
                      model = "McFL",
                      onlyCancer = FALSE,
                      finalTime = 50,
                      mu = 1e-4,
                      initSize = 5000,
                      keepPhylog = TRUE,
                      seed = NULL,
                      errorHitMaxTries = FALSE,
                      errorHitWallTime = FALSE)
plot(s1fd, show = "genotypes")
```

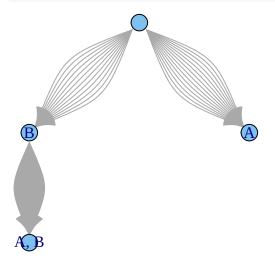




plotClonePhylog(s1fd, keepEvents = TRUE)



### plotClonePhylog(s2fd, keepEvents = TRUE)



Of course, it also works with absolute frequencies (code not executed for the sake of speed):

```
## Absolute
r5 <- data.frame(Genotype = c("WT", "A", "B", "A, B"),
                 Fitness = c("1",
                              "1.25 - .0025*(n_2)",
                              "1.25 - .0025*(n 1)",
                              "1.4"))
afe5 <- allFitnessEffects(genotFitness = r5,</pre>
                          frequencyDependentFitness = TRUE,
                          frequencyType = "abs")
set.seed(8)
s5 <- oncoSimulIndiv(afe5,
                      model = "McFL",
                      onlyCancer = FALSE,
                      finalTime = 100,
                      mu = 1e-4,
                      initSize = 5000,
                     keepPhylog = TRUE,
                      seed = NULL,
                      errorHitMaxTries = FALSE,
                      errorHitWallTime = FALSE)
plot(s5, show = "genotypes")
plot(s5, show = "genotypes", log = "y", type = "line")
mt <- allMutatorEffects(epistasis = c("A" = 0.1,</pre>
                                        "B" = 10)
set.seed(8)
s6 <- oncoSimulIndiv(afe5,
                      muEF = mt,
```

Note, though, that **evalAllGenotypesFitAndMut** does not currently work with frequency-dependent fitnes:

```
evalAllGenotypes(allFitnessEffects(genotFitness = r1fd,
                                     frequencyDependentFitness = TRUE,
                                     frequencyType = "rel",
                                     spPopSizes = c(10, 20, 30, 40)))
##
     Genotype Fitness
## 1
          WT
                   1.0
## 2
                   1.7
            \boldsymbol{A}
            B
## 3
                   1.6
        A, B
                   2.1
## 4
## Fitness is wrong
evalAllGenotypesFitAndMut(allFitnessEffects(genotFitness = r1fd,
                                     frequencyDependentFitness = TRUE,
                                     frequencyType = "rel",
                                     spPopSizes = c(10, 20, 30, 40)),
                     mtfd)
     Genotype Fitness MutatorFactor
##
## 1
            \boldsymbol{A}
                   1.4
                                   0.1
## 2
             В
                                  10.0
                   1.4
         A, B
                   1.6
                                   1.0
## 3
```

# 10.7 Can we use the BNB algorithm to model frequency-dependent fitness?

This question is similar to the one we address in 14.6. Briefly, the answer is yes. You can think of this as an approximation to an exact simulation of a stochastic system. You can also think of a delay in the system in the sense that the changes in rates due to changes in the frequencies of the different genotypes are updated at periodic

# 11 Additional examples of frequency-dependent fitness

In this section, we provide additional examples that use frequency-dependent fitness. As mentioned also in 10, **This functionality is only available in the freq-dep-fitness branch**. See: https://github.com/rdiaz02/OncoSimul/tree/freq-dep-fitness.

Note also that in most of these examples we make rather arbitrary and simple assumptions about the genetic basis of the different phenotypes or strategies (most are one-mutation-away from WT); see 10.4 and 10.2 (where we change the ancestor-dependent relationships). In some examples mutation rates are also very high, to speed up processes and because a high mutation rate is used as a procedure (a hack?) to quickly obtain descendants from WT (i.e., to get the game started with some representatives of the non-WT types).

Examples 11.1, 11.2, 11.3, 11.4, 11.5 were originally prepared by Sara Dorado Alfaro, Miguel Hernández del Valle, Álvaro Huertas García, Diego Mañanes Cayero, Alejandro Martín Muñoz as an exercisse for the course Programming and Statistics with R (Master's Degree in Bioinformatics and Computational Biology, Universidad Autónoma de Madrid), course 2019-20.

## 11.1 Rock-paper-scissors model in bacterial community

### 11.1.1 Introduction

This example is inspired by Kerr et al. (2002). It describes the relationship between three populations of *Escherichia coli*, that turns out to be very similar to a rock-paper-scissors game.

An E. coli community can have a specific strain of colicinogenic bacteria, that are capable of creating colicin, a toxin to which this special strain is resistant. The wild-type bacteria is killed by this toxin, but can mutate into a resistant strain.

So, there are three kinds of bacteria: wild-type (WT), colicinogenic (C) and resistant (R). The presence of C reduces the population of WT, but increases the population of R because R has an advantage over C, since R doesn't have the cost of creating the toxin. At the same time, WT has an advantage over R, because by losing the toxin receptors, R loses also some important functions. Therefore, every strain "wins" against one strain and "loses" against the other, creating a rock-paper-scissors game.

```
))
}
```

The equations are:

$$W(WT) = 1 + af_R - bf_C \tag{3}$$

$$W(C) = 1 + bf_{WT} - cf_R \tag{4}$$

$$W(R) = 1 + cf_C - af_{WT} \tag{5}$$

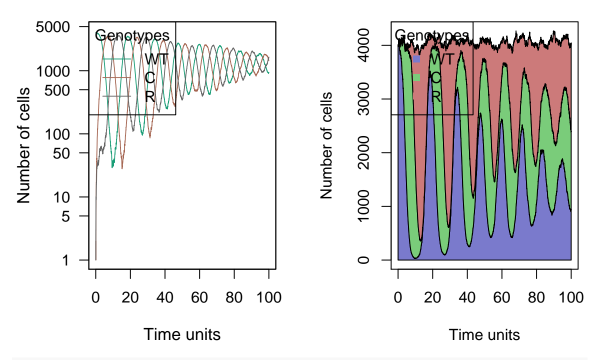
(6)

where  $f_{WT}$ ,  $f_C$  and  $f_R$  are the frequencies of WT, C and R, respectively.

#### 11.1.1.1 Case 1

We are going to study the scenario in which all the relationships have the same relative weight.

```
afcrs1 <- allFitnessEffects(genotFitness = crs(1, 1, 1),
                            frequencyDependentFitness = TRUE,
                            frequencyType = "rel")
resultscrs1 <- oncoSimulIndiv(afcrs1,
                              model = "McFL",
                              onlyCancer = FALSE,
                              finalTime = 100,
                              mu = 1e-2,
                              initSize = 4000,
                              keepPhylog = TRUE,
                              seed = NULL,
                              errorHitMaxTries = FALSE,
                              errorHitWallTime = FALSE)
op \leftarrow par(mfrow = c(1, 2))
plot(resultscrs1, show = "genotypes", type = "line", cex.lab=1.1,
     las = 1)
plot(resultscrs1, show = "genotypes", type = "stacked")
```



par(op)

An oscillatory equilibrium is reached, in which the same populations have a similar number of individuals but oscillates. This makes sense, because the rise on a particular strand will lead to a rise in the one that "wins" against it, and then to a rise in the one that "wins" against the second one, creating this cyclical behaviour. In the stacked plot we can see that the total population remains almost constant.

Note, though, that altering mutation rate (which is huge here) can change the results of the model.

#### 11.1.1.2 Case 2

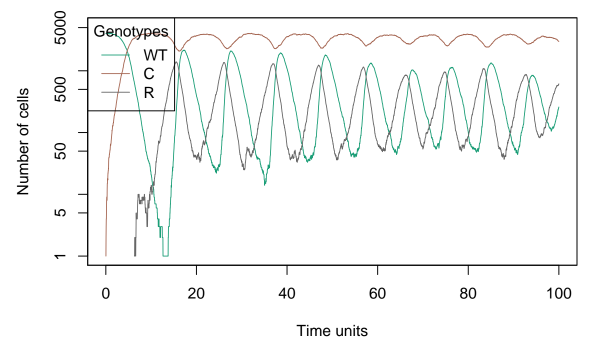
We are going to put a bigger weight in one of the coefficients, so a=10, b=1, c=1.

If we run multiple simulations, for example by doing

```
errorHitWallTime = FALSE)
```

we can verify there are two different scenarios.

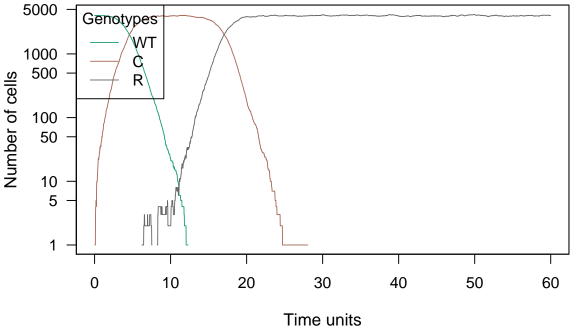
The first one is the one in which all the strains coexist, with the colicinogenic bacteria having a much bigger population.



In the second one, the wild type and the colicinogenic bacteria dissapear, so the resistant strain is the only one that survives.

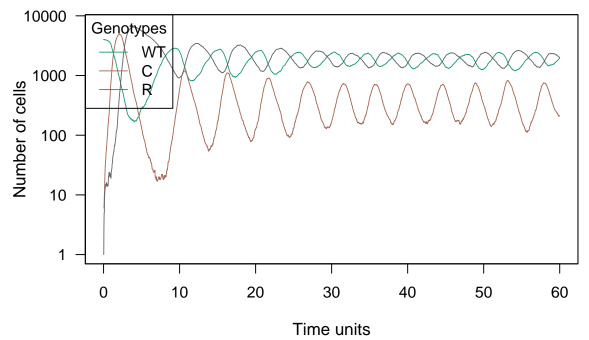
As above, though, decreasing the mutation rate can lead to a different solution and you will want to run the model for much longer to see the resistant strain appear and outcompete the others.

```
set.seed(3)
resultscrs2b <- oncoSimulIndiv(afcrs2,
                             model = "McFL",
                              onlyCancer = FALSE,
                              finalTime = 60,
                             mu = 1e-2,
                              initSize = 4000,
                             keepPhylog = TRUE,
                              seed = NULL,
                              errorHitMaxTries = FALSE,
                              errorHitWallTime = FALSE)
plot(resultscrs2b, show = "genotypes", type = "line", cex.lab=1.1,
     las = 1)
  5000
         Genotypes
                WT
```



#### 11.1.1.3 Case 3

Finally, we are going to put more weight in two coefficients, so a=1, b=5, c=5.



In all the cases all three strains survive, with C having a much smaller population than the other two.

# 11.2 Hawk and Dove example

The example we are going to show is one of the first games that Maynard Smith analyzed, for example in his classic Maynard Smith (1982) (see also, e.g., https://en.wikipedia.org/wiki/Chicken\_%28game%29).

In this game, the two competitors are subtypes of the same species but with different strategies. The Hawk first displays aggression, then escalates into a fight until it either wins or is injured (loses). The Dove first displays aggression, but if faced with major escalation runs for safety. If not faced with such escalation, the Dove attempts to share the resource (see the payoff matrix, for instance in https://en.wikipedia.org/wiki/Chicken\_%28game%29#Hawk%E2%80%93dove).

Given that the resource is given the value V, the damage from losing a fight is given cost C:

• If a Hawk meets a Dove he gets the full resource V to himself

- If a Hawk meets a Hawk half the time he wins, half the time he loses... so his average outcome is then V/2 minus C/2
- If a Dove meets a Hawk he will back off and get nothing 0
- If a Dove meets a Dove both share the resource and get V/2

The actual payoff however depends on the probability of meeting a Hawk or Dove, which in turn is a representation of the percentage of Hawks and Doves in the population when a particular contest takes place. That in turn is determined by the results of all of the previous contests. If the cost of losing C is greater than the value of winning V (the normal situation in the natural world) the mathematics ends in an stationary point (ESS), a mix of the two strategies where the population of Hawks is V/C.

In this case we assume a stable equilibrium in the population dynamics, that is, although there are external variations in the model, it recovers and returns to equilibrium.

We are going to simulate with OncoSimulR the situation in which the cost of losing C is greater than the value of gaining V ( $C=10,\,V=2$ ). We assume that both Hawk and Dove are derived from WT by one mutation (see also 10.4) and we will use very high mutation rates to get some hawks and doves from WT quickly (see above).

Before performing the simulation, let's look at the fitness of each competitor.

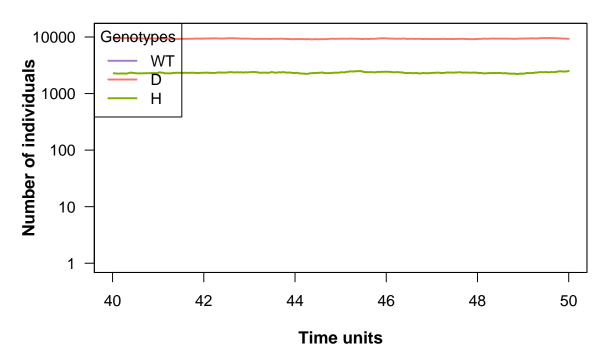
```
## Stablish Genotype-Fitnees mapping. D = Dove, H = Hawk
H D fitness <- function(c, v,</pre>
                    gt = c("WT", "H", "D")) {
  data.frame(Genotype = gt,
             Fitness = c(
               paste0("1"),
               paste0("1 + f_1 *", (v-c)/2, "+ f_2 *", v),
               paste0("1 + f_2 *", v/2)))
}
## Fitness Effects specification
HD competition <-allFitnessEffects(</pre>
  genotFitness = H_D_fitness(10, 2,
                         gt = c("WT", "H", "D")),
  frequencyDependentFitness = TRUE,
  frequencyType = "rel",
  spPopSizes = c(5000, 5000, 5000))
## Plot fitness landscape of genotype "H, D" evaluation
data.frame("Doves fitness" = evalGenotype(genotype = "D",
                                           fitnessEffects = HD_competition),
           "Hawks_fitness" = evalGenotype(genotype = "H",
                                           fitnessEffects = HD competition))
##
     Doves_fitness Hawks_fitness
```

```
## 1 1.333 0.3333
```

We observe that the penalty of fighting (C > V) benefits the dove in terms of fitness respect to the hawk.

```
## Simulated trajectories
## run only a few for the sake of speed
simulation <- oncoSimulPop(2,</pre>
                           mc.cores = 2,
                           HD competition,
                           model = "McFL", # There is no collapse
                           onlyCancer = FALSE,
                           finalTime = 50,
                           mu = 1e-2, # Quick emergence of D and H
                           initSize = 4000,
                           keepPhylog = TRUE,
                           seed = NULL,
                           errorHitMaxTries = FALSE,
                           errorHitWallTime = FALSE)
## Plot first trajectory as an example
plot(simulation[[1]], show = "genotypes", type = "line",
    xlim = c(40, 50),
     lwdClone = 2, ylab = "Number of individuals",
     main = "Hawk and Dove trajectory",
     col = c("#a37acc", "#f8776d", "#7daf00"),
     font.main=2, font.lab=2,
     cex.main=1.4, cex.lab=1.1,
     las = 1)
```

# Hawk and Dove trajectory



As mentioned above, mathematically when a stationary point (ESS) is reached the relative frequency of hawks is V/C and doves 1-(V/C). Considering f\_H as relative frequency of hawks and f\_D = 1-f\_H as frequency of doves:

Hawk:

$$1 + f_H * (v - c)/2 + (1 - f_H) * v$$

Dove:

$$1 + (1 - f_H) * v/2$$

Hawk = Dove:

$$1 + f_H * (v - c)/2 + (1 - f_H) * v = 1 + (1 - f_H) * v/2$$

Resolving for f\_H:

$$f_H = v/c$$

Therefore, the relative frequency of hawks in equilibrium is equal to V/C. In our case it would be 20% (C = 10, V = 2). Let's check it:

```
## Recover the final result from first simulation
result <- tail(simulation[[1]][[1]], 1)

## Get the number of organisms from each species
n_WT <- result[2]
n_D <- result[3]
n_H <- result[4]</pre>
```

To sum up, this example shows that when the risks of contest injury or death (the Cost C) is significantly greater than the potential reward (the benefit value V), the stable population will be mixed between aggressors and doves, and the proportion of doves will exceed that of the aggressors. This explains behaviours observed in nature.

# 11.3 Game Theory with social dilemmas of tumour acidity and vasculature

This example is based on Kaznatcheev et al. (2017). In this work, it is explained that the progression of cancer is marked by the acquisition of a number of hallmarks, including self-sufficiency of growth factor production for angiogenesis and reprogramming energy metabolism for aerobic glycolysis. Moreover, there is evidence of intra-tumour heterogeneity. Given that some cancer cells can not invest in something that benefits the whole tumor while others can free-ride on the benefits created by them (evolutionary social dilemmas), how do these population level traits evolve, and how are they maintained? The authors answer this question with a mathematical model that treats acid production through glycolysis as a tumour-wide public good that is coupled to the club good of oxygen from better vascularisation.

The cell types of the model are:

- VOP: VEGF (over)-producers.
- GLY: glycolytic cells.
- DEF: aerobic cells that do not call for more vasculature.

On the other hand, the micro-environmental parameters of the model are:

- a: the benefit per unit of acidification.
- v: the benefit from oxygen per unit of vascularisation.
- c: the cost of (over)-producing VEGF.

The fitness equations derived from those populations and parameters are:

$$W(GLY) = 1 + a * (f_1 + 1)$$
(7)

$$W(VOP) = 1 + a * f_1 + v * (f_2 + 1) - c$$
(8)

$$W(DEF) = 1 + a * f_1 + v * f_2$$
(9)

Where  $f_1$  is the GLY cells' frequency and  $f_2$  is the DEF cells' frequency at a given time. All fitness equations start from balance by the sum of 1.

Finally, depending of the parameter's values, the model can lead to three different situations (as in other examples, the different types are one mutation away from WT):

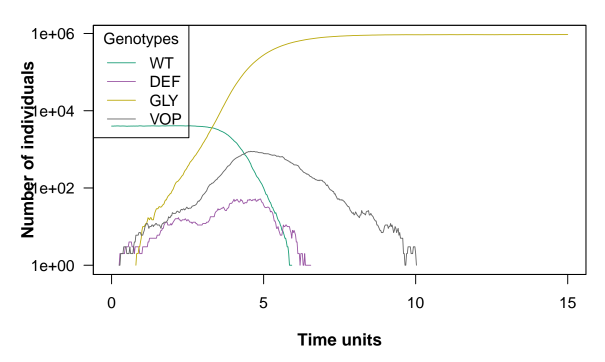
### 11.3.1 Fully glycolytic tumours:

If the fitness benefit of a single unit of acidification is higher than the maximum benefit from the club good for aerobic cells, then GLY cells will always have a strictly higher fitness than aerobic cells, and be selected for. In this scenario, the population will converge towards all GLY, regardless of the initial proportions (as long as there is at least some GLY in the population).

```
# Definition of the function for creating the corresponding dataframe.
avc <- function (a, v, c) {
  data.frame(Genotype = c("WT", "GLY", "VOP", "DEF"),
             Fitness = c("1",
                         paste0("1 + ",a," * (f 1 + 1)"),
                         paste0("1 + ",a," * f_1 + ",v," * (f_2 + 1) - ",c),
                         paste0("1 + ",a," * f 1 + ",v," * f 2")
                         ))
                          }
# Specification of the different effects on fitness.
afavc <- allFitnessEffects(genotFitness = avc(2.5, 2, 1),</pre>
                           frequencyDependentFitness = TRUE,
                           frequencyType = "rel")
## For real, you would probably want to run
## this multiple times with oncoSimulPop
simulation <- oncoSimulIndiv(afavc,</pre>
                           model = "McFL",
                            onlyCancer = FALSE,
                           finalTime = 15,
                           mu = 1e-3,
                            initSize = 4000,
                           keepPhylog = TRUE,
                            seed = NULL,
                            errorHitMaxTries = FALSE,
                            errorHitWallTime = FALSE)
```

```
# Representation of the plot of one simulation as an example (the others are
# highly similar).
plot(simulation, show = "genotypes", type = "line",
    ylab = "Number of individuals", main = "Fully glycolytic tumours",
    font.main=2, font.lab=2, cex.main=1.4, cex.lab=1.1, las = 1)
```

# **Fully glycolytic tumours**



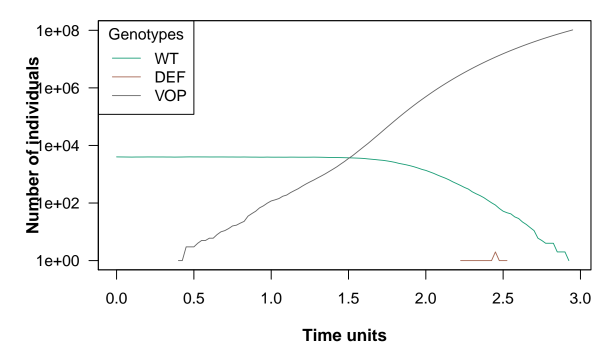
### 11.3.2 Fully angiogenic tumours:

If the benefit to VOP from their extra unit of vascularisation is higher than the cost c to produce that unit, then VOP will always have a strictly higher fitness than DEF, selecting the proportion of VOP cells towards 1. In addition, if the maximum possible benefit of the club good to aerobic cells is higher than the benefit of an extra unit of acidification, then for sufficiently high number of VOP, GLY will have lower fitness than aerobic cells. When both conditions are satisfied, the population will converge towards all VOP.

```
model = "McFL",
onlyCancer = FALSE,
finalTime = 15,
mu = 1e-4,
initSize = 4000,
keepPhylog = TRUE,
seed = NULL,
errorHitMaxTries = FALSE,
errorHitWallTime = FALSE)
```

```
## We get a huge number of VOP very quickly
## (too quickly?)
plot(simulation, show = "genotypes", type = "line",
    ylab = "Number of individuals", main = "Fully angiogenic tumours",
    font.main=2, font.lab=2, cex.main=1.4, cex.lab=1.1, las = 1)
```

# **Fully angiogenic tumours**

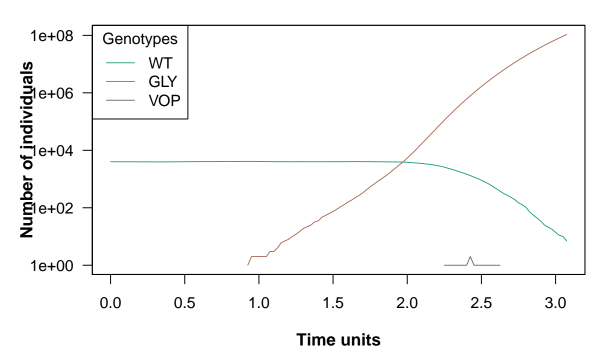


### 11.3.3 Heterogeneous tumours:

If the benefit from an extra unit of vascularisation in a fully aerobic group is lower than the cost c to produce that unit, then for a sufficiently low proportion of GLY and thus sufficiently large number of aerobic cells sharing the club good, DEF will have higher fitness than VOP. This will lead to a decrease in the proportion of VOP among aerobic cells and thus a decrease in the average fitness of aerobic cells. A lower fitness in aerobic cells will lead to an increase in the proportion of GLY until the aerobic groups (among which the club good is split) get sufficiently small and fitness starts to favour VOP over DEF, swinging the dynamics back.

```
# Definition of the function for creating the corresponding dataframe.
avc <- function (a, v, c) {
  data.frame(Genotype = c("WT", "GLY", "VOP", "DEF"),
             Fitness = c("1",
                         paste0("1 + ",a," * (f_1 + 1)"),
                         paste0("1 + ",a," * f_1 + ",v," * (f_2 + 1) - ",c),
                         paste0("1 + ",a," * f 1 + ",v," * f 2")
                         ))
                          }
# Specification of the different effects on fitness.
afavc <- allFitnessEffects(genotFitness = avc(7.5, 2, 1),</pre>
                           frequencyDependentFitness = TRUE,
                           frequencyType = "rel")
# Launching of the simulation (20 times).
simulation <- oncoSimulIndiv(afavc,</pre>
                           model = "McFL",
                           onlyCancer = FALSE,
                           finalTime = 25,
                           mu = 1e-4,
                           initSize = 4000,
                           keepPhylog = TRUE,
                           seed = NULL,
                            errorHitMaxTries = FALSE,
                           errorHitWallTime = FALSE)
# Representation of the plot of one simulation as an example (the others are
# highly similar).
plot(simulation, show = "genotypes", type = "line",
     ylab = "Number of individuals", main = "Heterogeneous tumours",
     font.main=2, font.lab=2, cex.main=1.4, cex.lab=1.1, las = 1)
```

# Heterogeneous tumours



### 11.4 Prostate cancer tumour–stroma interactions

This example is based on Basanta et al. (2012). The authors apply evolutionary game theory to model the behavior and progression of a prostate tumour formed by three different cell populations: stromal cells, a dependant tumour phenotype capable of co-opting stromal cells to support its growth and an independent tumour phenotype that does not require microenvironmental support, be it stromal associated or not. To enable this, the model has four variables, which is the minimum necessary to describe the relationships in terms of costs and benefits between the different types of cells and, of course, to describe the progression of the cancer.

The different cell types, hence, are as follows:

- 1. S: stromal cells.
- 2. D: microenvironmental-dependent tumour cells.
- 3. I: microenvironmental-independent tumour cells.

And the parameters that describe the relationships are as follows:

- $\alpha$ : benefit derived from the cooperation between a S cell and a D cell.
- $\beta$ : cost of extracting resources from the microenvironment.
- $\gamma$ : cost of being microenvironmentally independent.
- $\rho$ : benefit derived by D from paracrine growth factors produced by I cells.

Table 18 shows the payoffs for each cell type when interacting with others. We consider no other phenotypes are relevant in the context of the game and disregard spatial considerations.

Table 18: Payoff table that represents the interactions between the three cell types considered by de model

	S	D	I
$\overline{\mathbf{S}}$	0	$\alpha$	0
D	$1 + \alpha - \beta$	$1-2\beta$	$1 - \beta + \rho$
Ι	$1-\gamma$	$1 - \gamma$	$1-\gamma$

As is@basanta investigating 2012, the I cells are relatively independent from the microenvironment and produce their own growth factors (e.g. testosterone) and thus are considered to have a comparatively constant fitness  $(1-\gamma)$ , where  $\gamma$ represents the fitness cost for I cells to be independent. The D cells rely more on their microenvironment for survival and growth at a fitness cost  $(\beta)$  that represents the scarcity of resources or space that I cells can procure themselves. A resource-poor microenvironment would then be characterised by a higher value of  $\beta$ . As I cells produce space and shareable growth factors, this model assumes that D cells derive a fitness advantage from their interactions with I cells represented by the variable  $\rho$ . On the other hand, D cells interacting with other D cells will have a harder time sharing existing microenvironmental resources with other equally dependant cells and thus are assumed to have double the cost  $2\beta$  for relying on the microenvironment for survival and growth and thus have a fitness of  $1^{\circ}2\beta$ . The S cells can interact with tumour cells. In a normal situation, this population are relatively growth quiescent with low rates of proliferation and death. For this reason the fitness benefit derived by stromal cells from the interactions with tumour cells is assumed to be zero. However, they are able to undergo rapid proliferation and produce growth factors if they are stimulated by factors produced by I cells, giving rise to a mutualistic relationship. This relationship is represented by the parameter  $\alpha$ . A low  $\alpha$  represents tumours in which the stroma cannot be co-opted and vice versa.

From these variables, the fitness of each cell population (W(S), W(I), W(D)) is as follows:

$$W(S) = 1 + f_3 \alpha \tag{10}$$

$$W(I) = 2 - \gamma \tag{11}$$

$$W(D) = 1 + (1 - f_2 - f_3)(1 - \beta + \alpha) + f_2(1 - \beta + \rho) + f_3(1 - 2\beta) + 1 - \beta + \alpha + f_2(\rho - \alpha) - f_3(\beta + \alpha)$$
(12)

where  $f_2$  is the frequency of I cells and  $f_3$  is the frequency of D cells at a given time. All fitness equations start from balance by the sum of 1.

### 11.4.1 Simulations

First, we define the fitness of the different genotypes (see Equations (10), (11) and (12)) through the function *fitness\_rel* that builds a data frame.

It is important to note that this program models a situation where, from a WT cell population, the rest of the cell population types are formed. However, this model has also stromal cells that are not formed from a WT, since they are not tumour cells although interacting with it. Hence, for this model, we can not represent scenarios with total biological accuracy, something that we must consider when interpreting the results.

Then, we are going to model different scenarios that represent different biological situations. In this case, we are going to explain four possible situations.

**Note:** for these simulations the values of paratemers are normalised in the range (0 : 1) so 1 represents the maximum value for any parameter being positive of negative to fitness depending on the parameter.

### 11.4.1.1 First scenario

In this simulation, we are modelling a situation where the environment is relatively resource-poor. In addition, we set a intermediate cooperation between D-D and D-I and a very low benefit from coexistence of D with I.

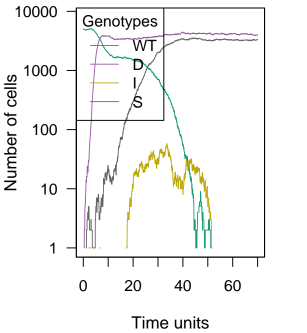
- $\alpha$  (a) = 0.5: intermediate cooperation between D and D cells.
- $\beta$  (b) = 0.7: relatively resource-poor microenvironment.
- $\rho$  (p) = 0.1: low benefit of D cells. -  $\gamma$  (g) = 0.8: high cost of independence of I cells.

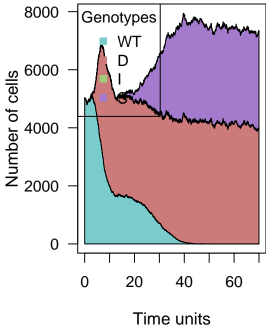
We can observe that high values of  $\alpha$  and low values of  $\rho$  are translated in a larger profit of D cells from his interaction with S cells than from his interaction with I cells. Also, because of the high cost of independence of I cells  $(\gamma)$ , it is not surprise that this population ends up becoming extinct. Finally, the tumour is composed by two cellular types: D and S cells.

```
scen1 <- allFitnessEffects(genotFitness = fitness_rel(a = 0.5, b = 0.7,</pre>
                                                        r = 0.1, g = 0.8),
                           frequencyDependentFitness = TRUE,
                           frequencyType = "rel")
set.seed(1)
simulScen1 <- oncoSimulIndiv(scen1,</pre>
                                model = "McFL",
                                onlyCancer = FALSE,
                                finalTime = 70,
                                mu = 1e-4,
                                initSize = 5000,
                                keepPhylog = TRUE,
                                seed = NULL,
                                errorHitMaxTries = FALSE,
                                errorHitWallTime = FALSE)
op \leftarrow par(mfrow = c(1, 2))
plot(simulScen1, show = "genotypes", type = "line",
     main = "First scenario",
     cex.main = 1.4, cex.lab = 1.1,
     las = 1)
plot(simulScen1, show = "genotypes",
     main = "First scenario",
     cex.main = 1.4, cex.lab = 1.1,
     las = 1)
```

## First scenario

# First scenario





```
par(op)
```

To understand the stability of the results, we should run multiple simulations. We will not pursue that here. Note that the results can be sensitive to the initial population size and the mutation rate.

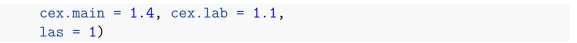
### 11.4.1.2 Second scenario

In this case, we set  $\alpha$  lower than in the first scenario and we enable the independence of I cells through a lower  $\gamma$ .

- $\alpha$  (a) = 0.3: low cooperation between D cells.
- $\beta$  (b) = 0.7: relatively resource-poor microenvironment.
- $\rho$  (r) = 0.1: low benefit of D cells from coexisting with I cells.
- $\gamma$  (g) = 0.7: lower cost of independence of I cells than in the first scenario.

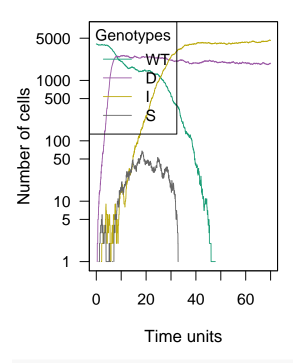
Because of we are easing the possibility of independence of I cells, instead of extinguishing as in the first scenario, they compose the bulk of the tumour along with D cells in spite of the low benefit of cooperation between them (low  $\rho$ ). Besides, we can observe that the population of I cells is bigger than the population of D cells, being at the end of the simulation in balance. On the other hand, stromal cells drop at the beginning of the simulation.

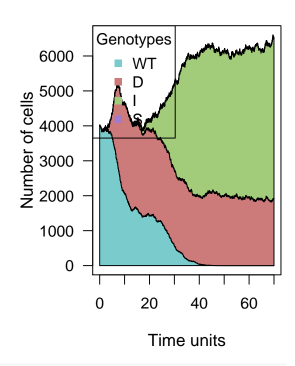
```
scen2 <- allFitnessEffects(genotFitness = fitness_rel(a = 0.3, b = 0.7,</pre>
                                                       r = 0.1, g = 0.7),
                           frequencyDependentFitness = TRUE,
                           frequencyType = "rel")
set.seed(1)
simulScen2 <- oncoSimulIndiv(scen2,
                               model = "McFL",
                               onlyCancer = FALSE,
                               finalTime = 70,
                               mu = 1e-4,
                               initSize = 4000,
                               keepPhylog = TRUE,
                               seed = NULL,
                               errorHitMaxTries = FALSE,
                               errorHitWallTime = FALSE)
op \leftarrow par(mfrow = c(1, 2))
plot(simulScen2, show = "genotypes", type = "line",
     main = "Second scenario",
     cex.main = 1.4, cex.lab = 1.1,
     las = 1)
plot(simulScen2, show = "genotypes",
     main = "Second scenario",
```



## Second scenario

# **Second scenario**





par(op)

### 11.4.1.3 Third scenario

In this case, we have a extreme situation where the microenvironment is rich (high  $\beta$ ) and the independence costs are very low ( $\gamma$ ) in relation with the previous scenarios.

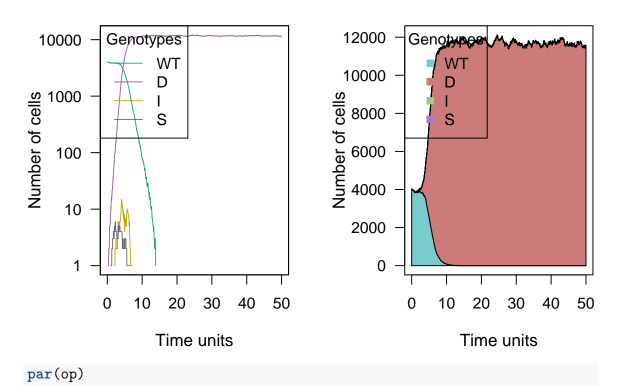
- $\alpha$  (a) = 0.2: low cooperation between D cells.
- $\beta$  (b) = 0.3: rich microenvironment, being beneficial for D cells.
- $\rho$  (r) = 0.1: low benefit of D cells from coexisting with I cells.
- $\gamma$  (g) = 0.3: independence costs very low, being beneficial for I cells.

Although  $\gamma$  and  $\rho$  are very low, which could make us think that I cells will control the tumour, we can observe that the fact that the microenvironment is very rich (with a low value of  $\beta$ ) allows to D cells lead the progression of the tumour over the rest of cell populations, including I cells.

```
onlyCancer = FALSE,
                                finalTime = 50,
                               mu = 1e-4,
                                initSize = 4000,
                               keepPhylog = TRUE,
                                seed = NULL,
                                errorHitMaxTries = FALSE,
                                errorHitWallTime = FALSE)
op \leftarrow par(mfrow = c(1, 2))
plot(simulScen3, show = "genotypes", type = "line",
     main = "Third scenario",
     cex.main = 1.4, cex.lab = 1.1,
     las = 1)
plot(simulScen3, show = "genotypes",
     main = "Third scenario",
     cex.main = 1.4, cex.lab = 1.1,
     las = 1)
```

## Third scenario

## Third scenario



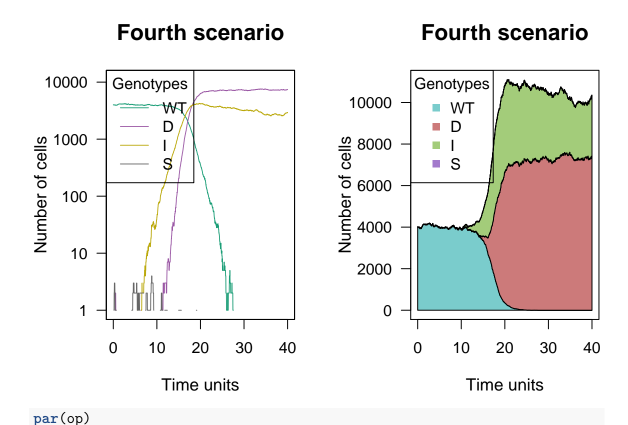
### 11.4.1.4 Fourth scenario

This is a variation of the third scenario to illustrate that, if we set a microenvironment more rich than in the previous scenario, we get a cooperation between D and I cells, although we can still observe the superiority of D cells over I cells.

•  $\alpha$  (a) = 0.2: low cooperation between D cells.

- $\beta$  (b) = 0.4: lower richness than in the previous scenario.
- $\rho$  (r) = 0.1: low benefit of D cells from coexisting with I cells.
- $\gamma$  (g) = 0.3: independence costs very low, being beneficial for I cells.

```
scen4 <- allFitnessEffects(genotFitness = fitness_rel(a = 0.2, b = 0.4,</pre>
                                                       r = 0.1, g = 0.3),
                           frequencyDependentFitness = TRUE,
                           frequencyType = "rel")
## Set a different seed to show the results better since
## with set.seed(1) the progression of I cells was not shown
set.seed(2)
simulScen4 <- oncoSimulIndiv(scen4,</pre>
                               model = "McFL",
                               onlyCancer = FALSE,
                               finalTime = 40,
                               mu = 1e-4,
                               initSize = 4000,
                               keepPhylog = TRUE,
                               seed = NULL,
                               errorHitMaxTries = FALSE,
                               errorHitWallTime = FALSE)
op \leftarrow par(mfrow = c(1, 2))
plot(simulScen4, show = "genotypes", type = "line",
     main = "Fourth scenario",
     cex.main = 1.4, cex.lab = 1.1,
     las = 1)
plot(simulScen4, show = "genotypes",
     main = "Fourth scenario",
     cex.main = 1.4, cex.lab = 1.1,
     las = 1)
```



In this case, we can see that there is more variation in the size of population of I cells. There are cases where the I cell population cooperates with D cells and, in others, there is not cooperation. You can examine this running multiple simulations (or manually rerun the example above changing the seed).

# 11.5 Evolutionary Dynamics of Tumor-Stroma Interactions in Multiple Myeloma

This example is based on Sartakhti et al. (2016). The authors provide a frequency-dependent model to study the growth of malignant plasma cells in multiple myeloma. Assuming that cancer cells and stromal cells cooperate by exchanging diffusible factors, the study is carried out in the framework of evolutionary game theory.

We first need to define a payoff strategy for this kind of scenario. The following definitions are needed:

- There are n phenotypes in a population denoted by  $\{P_1, \ldots, P_n\}$ .
- Each phenotype can produce one diffusible factor  $\{G_1, \ldots, G_n\}$ .
- Each diffusible factor j has a different effect  $r_{i,j}$  on the other phenotypes i.
- The cost for  $P_i$  for growth factor  $G_i$  is denoted as  $c_i$ .
- *M* is the number of cells within the diffusion range.
- There are  $M_j$  individuals of type  $P_j$  among the other group members.

Then, the payoff for strategy  $P_i$  is:

$$\pi_{P_j}(M_1, \dots, M_n) = \frac{(M_j + 1) \times c_j}{M} r_{j,j} + \sum_{i=1, i \neq j}^n \frac{M_i \times c_i}{M} r_{j,i} - c_j . \tag{13}$$

In multiple Myeloma we have three different types of cells that have autocrine and paracrine effects on the cells within their diffusion range: Malignant plasma cells (MM), Osteoblasts (OB) and Osteoclasts (OC). The specification of fitness is the following (see [10]):

$$W_{OC} = \frac{(f_1(M-1)+1)c_1}{M}r_{11} + \frac{((1-f_3)(M-1)-f_1(M-1)-1)c_2}{M}r_{12}$$
 (14)

$$+\frac{(M-(1-f_3)(M-1))c_3}{M}r_{13}-c_1\tag{15}$$

$$W_{OB} = \frac{(f_2(M-1)+1)c_2}{M}r_{22} + \frac{((1-f_1)(M-1)-f_2(M-1)-1)c_3}{M}r_{23}$$
 (16)

$$+\frac{(M-(1-f_1)(M-1))c_1}{M}r^{21}-c_2\tag{17}$$

$$W_{MM} = \frac{(f_3(M-1)+1)c_3}{M}r33 + \frac{((1-f_2)(M-1)-f_3(M-1)-1)c_1}{M}r31 \quad (18)$$

$$+\frac{(M-(1-f_2)(M-1))c_2}{M}r32-c_3, \qquad (19)$$

where  $f_1$ ,  $f_2$  and  $f_3$  denote the frequency of the phenotype OC, OB and MM in the population. The multiplication factors for diffusible factors produced by the cells are shown in the following table (taken from Sartakhti et al. (2016)):

{r smyeloma, eval=TRUE, echo=FALSE, fig.cap="Multiplication factorin myeloma interaction. Table 1 in Sartakhti et al., 2016, 'Evolutionary Dynamics of Tumor-Stroma Interactions in Multiple Myeloma' https://doi.org/10.1371/."} knitr::include\_graphics("Myeloma\_interaction.png")

Several scenarios varying the values of the parameters are shown in Sartakhti et al. (2016). Here we reproduce some of them.

### 11.5.1 Simulations

First, we define the fitness of the different genotypes through the function fitness\_rel.

It is important to note that, in order to exactly reproduce the experiments of the paper, we need to create an initial population with three different types of cell, but we do not need the presence of a wild type. For this reason, we will increase the probability of mutation of the wild type, which will disappear in early stages of the simulation; this is a procedure we have used before in several cases too (e.g., 11.2). This is something that we must consider when interpreting the results.

### 11.5.2 Scenario 1

Here we model a common situation in multiple myeloma in which  $c_1 < c_2 < c_3$ . In the presence of a small number of MM cells, the stable point on the OB-OC border becomes a saddle point and clonal selection leads to a stable coexistence of OC and MM cells. Parameters for the simulation can be seen in the R code.

```
N <- 40000

M <- 10

c1 <- 1

c2 <- 1.2

c3 <- 1.4

r11 <- 0

r12 <- 1

r13 <- 2.5

r21 <- 1

r22 <- 0

r23 <- -0.3

r31 <- 2.5

r32 <- 0

r33 <- 0
```

```
fe_cells <-
    allFitnessEffects(
         genotFitness =
             f_cells(c1, c2, c3, r11, r12, r13,
                      r21, r22, r23, r31, r32, r33, M,
                      gt = c("WT", "OC", "OB", "MM")),
         frequencyDependentFitness = TRUE,
         frequencyType = "rel")
## All single-gene genotypes as input to to_genotFitness_std
## Simulated trajectories
set.seed(2)
simulation <- oncoSimulIndiv(fe_cells,</pre>
                             model = "McFL",
                              onlyCancer = FALSE,
                             finalTime = 30,
                             mu = c("OC"=1e-1, "OB"=1e-1, "MM"=1e-4),
                              initSize = N,
                             keepPhylog = FALSE,
                              seed = NULL,
                              errorHitMaxTries = FALSE,
                              errorHitWallTime = FALSE)
                                            #Plot trajectorie
op \leftarrow par(mfrow = c(1, 2))
plot(simulation, show = "genotypes", type = "line")
plot(simulation, show = "genotypes")
           Genotypes
                                                    Genotypes
                 ŴΤ
                                                        WT
                  MM
                                                        MM
                                              30000
                  ÓB
                                                        OB
Number of cells
                                         Number of cells
     1000
                  OC
                                                        OC
     100
                                              10000
     10
                                              0
          0
              5
                  10
                         20
                                                   0
                                                       5
                                                          10
                                                                  20
                                 30
                                                                          30
                  Time units
                                                           Time units
```

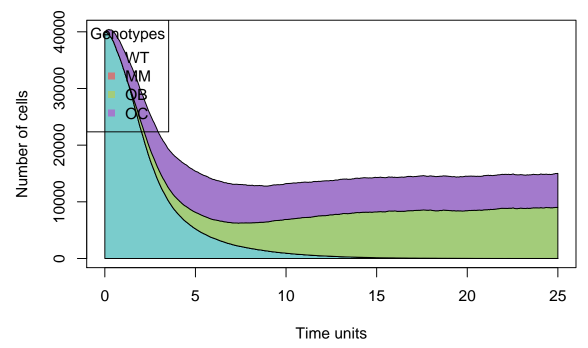
```
par(op)
```

Clearly, the appearance of MM cells quickly brings the system to an equilibrium point, which is stable. OB cells are extinct and cancer has propelled.

#### 11.5.3 Scenario 2

As in the second scenario of [10] (c1 = c2 = c3) configuration A (upper row in the grid of images). We should find one stable point on the OC-OB edge under certain conditions, which are met in the example. Further information about the parameters can be found in the R code shown below.

```
N < -40000
M < -10
c1 <- 1
c2 <- 1
c3 <- 1
r11 <- 0
r12 <- 1
r13 < -0.5
r21 <- 1
r22 <- 0
r23 < -0.3
r31 < -0.5
r32 <- 0
r33 <- 0
fe cells <-
    allFitnessEffects(
        genotFitness =
            f_cells(c1, c2, c3, r11, r12, r13,
                    r21, r22, r23, r31, r32, r33, M,
                    gt = c("WT", "OC", "OB", "MM")),
        frequencyDependentFitness = TRUE,
        frequencyType = "rel")
## All single-gene genotypes as input to to_genotFitness_std
## Simulated trajectories
set.seed(1)
simulation <- oncoSimulIndiv(fe_cells,</pre>
                            model = "McFL",
                            onlyCancer = FALSE,
                            finalTime = 25,
                            mu = c("OC"=1e-1, "OB"=1e-1, "MM"=1e-4),
                            initSize = N,
                            keepPhylog = FALSE,
                            seed = NULL,
```



As expected, under these conditions, MM cells are not able to propagate. The equilibrium point in the OB-OC edge is stable, resisting small variations in the number of MM cells.

# Measures of evolutionary predictability and genotype diversity

Several measures of evolutionary predictability have been proposed in the literature (see, e.g., Szendro, Franke, et al. (2013) and references therein). We provide two, Lines of Descent (LOD) and Path of the Maximum (POM), following Szendro, Franke, et al. (2013); we also provide a simple measure of diversity of the actual genotypes sampled.

In Szendro, Franke, et al. (2013) "(...) paths defined as the time ordered sets of genotypes that at some time contain the largest subpopulation" are called "Path of the Maximum" (POM) (see their p. 572). In our case, POM are obtained by finding the clone with largest population size whenever we sample and, thus, the POMs will be affected by how often we sample (argument sampleEvery), since we are running a continuous time process.

Szendro, Franke, et al. (2013) also define Lines of Descent (LODs) which "(...) represent the lineages that arrive at the most populated genotype at the final time". In that same page (572) they provide the details on how the LODs are obtained. Starting with version 2.9.2 of OncoSimulR I only provide an implementation where a single LOD per simulation is returned, with the same meaning as in Szendro, Franke, et al. (2013).

To briefly show some output, we will use again the 5.5 example.

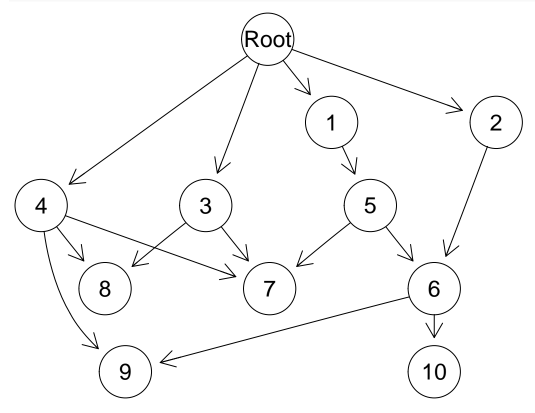
```
pancr <- allFitnessEffects(</pre>
    data.frame(parent = c("Root", rep("KRAS", 4), "SMAD4", "CDNK2A",
                           "TP53", "TP53", "MLL3"),
               child = c("KRAS", "SMAD4", "CDNK2A",
                          "TP53", "MLL3",
                          rep("PXDN", 3), rep("TGFBR2", 2)),
               s = 0.05, sh = -0.3, typeDep = "MN"))
pancr16 <- oncoSimulPop(16, pancr, model = "Exp",</pre>
                        mc.cores = 2)
## Look a the first POM
str(POM(pancr16)[1:3])
## List of 3
## $ : chr [1:2] "" "KRAS"
## $ : chr [1:2] "" "KRAS"
## $ : chr [1:3] "" "KRAS" "KRAS, MLL3, TP53"
LOD(pancr16)[1:2]
## [[1]]
## [17 ""
               "KRAS"
##
## [[2]]
```

```
## [1] "" "KRAS"
## The diversity of LOD (lod_single) and POM might or might not
## be identical
diversityPOM(POM(pancr16))
## [1] 1.124
diversityLOD(LOD(pancr16))
## [1] 0.6886
## Show the genotypes and their diversity (which might, or might
## not, differ from the diversity of LOD and POM)
sampledGenotypes(samplePop(pancr16))
##
## Subjects by Genes matrix of 16 subjects and 7 genes.
            Genotype Freq
## 1
       CDNK2A, KRAS
                       1
## 2
                KRAS
                       13
## 3
          KRAS, MLL3 1
## 4 KRAS, MLL3, TP53
                       1
##
## Shannon's diversity (entropy) of sampled genotypes: 0.6886
```

# 13 Generating random DAGs for restrictions

You might want to randomly generate DAGs like those often found in the literature on Oncogenetic trees et al. Function simOGraph might help here.

```
## No seed fixed, so reruns will give different DAGs.
(a1 <- simOGraph(10))
       Root 1 2 3 4 5 6 7 8 9 10
##
         0 1 1 1 1 0 0 0 0 0
## Root
         0000010000
## 1
## 2
         0000001000
         0000000110
## 3
         0 0 0 0 0 0 0 1 1 1
## 4
         0 0 0 0 0 0 1 1 0 0
## 5
## 6
         0000000001
         00000000000
## 8
         00000000000
## 9
         00000000000
         00000000000
## 10
library(graph) ## for simple plotting
plot(as(a1, "graphNEL"))
```



Once you obtain the adjacency matrices, it is for now up to you to convert them into appropriate posets or fitnessEffects objects.

Why this function? I searched for, and could not find any that did what I wanted, in particular bounding the number of parents, being able to specify the approximate

depth<sup>15</sup> of the graph, and optionally being able to have DAGs where no node is connected to another both directly (an edge between the two) and indirectly (there is a path between the two through other nodes). So I wrote my own code. The code is fairly simple to understand (all in file generate-random-trees.R). I would not be surprised if this way of generating random graphs has been proposed and named before; please let me know, best if with a reference.

Should we remove direct connections if there are indirect? Or, should we set removeDirectIndirect = TRUE? Setting removeDirectIndirect = TRUE is basically asking for the transitive reduction of the generated DAG. Except for Farahani and Lagergren (2013) and Ramazzotti et al. (2015), none of the DAGs I've seen in the context of CBNs, Oncogenetic trees, etc, include both direct and indirect connections between nodes. If these exist, reasoning about the model can be harder. For example, with CBN (AND or CMPN or monotone relationships) adding a direct connection makes no difference iff we assume that the relationships encoded in the DAG are fully respected (e.g., all  $s_h = -\infty$ ). But it can make a difference if we allow for deviations from the monotonicity, specially if we only check for the satisfaction of the presence of the immediate ancestors. And things get even trickier if we combine XOR with AND. Thus, I strongly suggest you leave the default removeDirectIndirect = TRUE. If you change it, you should double check that the fitnesses of the possible genotypes are what you expect. In fact, I would suggest that, to be sure you get what you think you should get, you convert the fitness from the DAG to a fitness table, and pass that to the simulations, and this requires using non-exposed user functions; to give you an idea, this could work (but you've been warned: this is dangerous!)

<sup>&</sup>lt;sup>15</sup>Where depth is defined in the usual way to mean smallest number of nodes —or edges— to traverse to get from the bottom to the top of the DAG.

# 14 FAQ, odds and ends

# 14.1 What we mean by "clone"; and "I want clones disregarding passengers"

In this vignette we often use "clone" or "genotype" interchangeably. A clone denotes a set of cells that have identical genotypes. So if you are using a fitness specification with four genes (i.e., your genome has only four loci), there can be up to  $16 = 2^4$  different genotypes or clones. Any two entities that differ in the genotype are different clones. And this applies regardless of whether or not you declare that some genes (loci) are drivers or not. So if you have four genes, it does not matter whether only the first or all four are regarded as drivers; you will always have at most 16 different clones or 16 different genotypes. Of course you can arrive at the same clone/genotype by different routes. Just think about loci A and B in our four-loci genome, and how you can end up with a cell with both A and B mutated.

Analogously, if you have 100 genes, 10 drivers and 90 passengers, you can have up to  $2^{100}$  different clones or genotypes. Sure, one cell might have driver A mutated and passenger B mutated, and another cell might have driver A mutated and passenger C mutated. So if you only look at drivers you might be tempted to say that they are "the same clone for all practical purposes"; but they really are not the same clone as they differ in their genotype and this makes a lot of difference computationally.

If you want summaries of simulations that collapse over some genes (say, some "passengers", the 90 passengers we just mentioned) look at the help for samplePop, argument geneNames. This would allow you, for instance, to look at the diversity of clones/genotypes, considering as identical those genotypes that only differ in genes you deem relevant; something similar to defining a "drivers' clone" as the set formed from the union of all sets of cells that have identical genotype with respect to only the drivers (so that in the example of "A, B" and "A, C" just mentioned both cells would be considered "the same clone" as they only differ with respect to passengers). However, this "disregard some genes" only applies to summaries of simulations once we are done simulating data. OncoSimulR will always track clones, as defined above, regardless of whether many of those clones have the same genotype if you were to only focus on driver genes; see also section 14.2.

Labeling something as a "driver", therefore, does not affect what we mean by clone. Yes, labeling something as a driver can affect when you stop simulations if you use detectionDrivers as a stopping mechanism (see section 6.3). But, again, this has nothing to do with the definition of "clone".

If this is all obvious to you, ignore it. I am adding it here because I've seen strange misunderstandings that eventually could be traced to the apparently multiple meanings of clone. (And to make the story complete, Mather, Hasty, and Tsimring (2012) use the expression "class" —e.g., Algorithm 4 in the paper, Algorithm 5 in the supplementary material).

# 14.2 Does OncoSimulR keep track of individuals or of clones? And how can it keep track of such large populations?

OncoSimulR keeps track of clones, where a clone is a set of cells that are genetically identical (note that this means completely identical over the whole set of genes/markers you are using; see section 14.1). We do not need to keep track of individual cells because, for all purposes, and since we do not consider spatial structure, two or more cells that are genetically identical are interchangeable. This means, for instance, that the computational cost of keeping a population of a single clone with 1 individual or with  $10^9$  individuals is exactly the same: we just keep track of the genotype and the number of cells. (Sure, it is much more likely we will see a mutation soon in a clone with  $10^9$  cells than in a clone with 1, but that is a different issue.)

Of course, the entities that die, reproduce, and mutate are individual cells. This is of course dealt with by tracking clones (as is clearly shown by Algorithms 4 and 5 in Mather, Hasty, and Tsimring (2012)). Tracking individuals, as individuals, would provide no advantage, but would increase the computational burden by many orders of magnitude.

### 14.2.1 sampleEvery, keepPhylog, and pruning

At each sampling time (where sampleEvery determines the time units between sampling times) the abundance of all the clones with number of cells > 0 is recorded. This is the structure that at the end of the run is converted into the pops.by.time matrix.

Now, some clones might arise from mutation between successive population samples but these clones might be extinct by the time we take a population sample. These clones do not appear in the pops.by.time matrix because, as we just said, they have 0 cells at the time of sampling. Of course, some of these clones might appear again later and reach a size larger than 0 at some posterior sampling time; it is at this time when this/these clone(s) will appear in the pops.by.time matrix. This pruning of clones with 0 cells can allow considerable savings in computing time (OncoSimulR needs to track the genotype of clones, their population sizes, their birth, death, and mutation rates, their next mutation time and the last time they were updated and thus it is important that we only loop over structures with information that is really needed).

However, we still need to track clones as clones, not simply as classes such as "number of mutated genes". Therefore, very large genomes can represent a problem if they lead to the creation and tracking of many different clones (even if they have the same number of mutated genes), as we have seen, for instance, in section 2.3. In this case, programs that only keep track of numbers of mutated genes or of drivers, not individual clones, can of course achieve better speed.

What about the genealogy? If you ask OncoSimulR to keep track of the complete parent-child relationships (keepPhylog = TRUE), you might see in the genealogy

clones that are not present in pops.by.time if these are clones that never had a population size larger than 0 at any sampling time. To give an example, suppose that we will take population samples at times 0, 1, and 2. Clone A, with a population size larger than 0 at time 1, gives rise at time 1.5 to clone B; clone B then gives rise to clone C at time 1.8. Finally, suppose that at time 2 only clone C is alive. In other words, when we carry out the update of the population with Algorithm 5 from Mather, Hasty, and Tsimring (2012), clones A and B have size 0. Now, at time 1 clones B and C did not yet exist, and clone B is never alive at times 1 or 2. Thus, clone B is not present in pops.by.time. But we cannot remove clone B from our genealogy if we want to reflect the complete genealogy of C. Thus, pops.by.time will show only clones A and C (not B) but the complete genealogy will show clones A, B, C (and will show that B appeared from A at time 1.5 and C appeared from B at time 1.8). Since function plotClonePhylog offers a lot of flexibility with respect to what clones to show depending on their population sizes at different times, you can prevent being shown B, but its existence is there should you need it (see also 2.3.5).

### 14.3 Dealing with errors in "oncoSimulPop"

When running OncoSimulR under Windows mclapply does not use multiple cores, and errors from oncoSimulPop are reported directly. For example:

Under POSIX operating systems (e.g., GNU/Linux or Mac OSX) oncoSimulPop can ran parallelized by calling mclapply. Now, suppose you did something like

The warning you are seeing tells you there was an error in the functions called by mclapply. If you check the help for mclapply you'll see that it returns a try-error object, so we can inspect it. For instance, we could do:

### pancrError[[1]]

But the output of this call might be easier to read:

### pancrError[[1]][1]

And from here you could see the error that was returned by oncoSimulIndiv: initSize < 1 (which is indeed true: we pass initSize = 1e-5).

# 14.4 Whole tumor sampling, genotypes, and allele counts: what gives? And what about order?

You are obtaining genotypes, regardless of order. When we use "whole tumor sampling", it is the frequency of the mutations in each gene that counts, not the order. So, for instance, "c, d" and "c, d" both contribute to the counts of "c" and "d". Similarly, when we use single cell sampling, we obtain a genotype defined in terms of mutations, but there might be multiple orders that give this genotype. For example, d > c and c > d both give you a genotype with "c" and "d" mutated, and thus in the output you can have two columns with both genes mutated.

# 14.5 Doesn't the BNB algorithm require small mutation rates for it to be advantageous?

As discussed in the original paper by Mather, Hasty, and Tsimring (2012) (see also their supplementary material), the BNB algorithm can achieve considerable speed advantages relative to other algorithms especially when mutation events are rare relative to birth and death events; the larger the mutation rate, the smaller the gains compared to other algorithms. As mentioned in their supplementary material (see p.5) "Note that the 'cost' of each step in BNB is somewhat higher than in SSA [SSA is the original Gillespie's Stochastic Simulation Algorithm] since it requires generation of several random numbers as compared to only two uniform random numbers for SSA. However this cost increase is small compared with significant benefits of jumping over birth and death reactions for the case of rare mutations."

Since the earliest versions, OncoSimulR has provided information to assess these issues. The output of function oncoSimulIndiv includes a list called "other" that itself includes two lists named "minDMratio" and "minBMratio", the smallest ratio, over all simulations, of death rate to mutation rate or birth rate to mutation rate, respectively. As explained above, the BNB algorithm thrives when those are large. Note, though, we say "it thrives": these ratios being large is not required for the BNB algorithm to be an exact simulation algorithm; these ratios being large make BNB comparatively much faster than other algorithms.

# 14.6 Can we use the BNB algorithm with state-dependent birth or death rates?

As discussed in the original paper by Mather, Hasty, and Tsimring (2012) (see sections 2.6 and 3.2 of the paper and section E of the supplementary material), the BNB algorithm can be used as an approximate stochastic simulation algorithm "(...) with non-constant birth, death, and mutation rates by evolving the system with a BNB step restricted to a short duration t." (p. 9 in supplementary material). The justification is that "(...) the propensities for reactions can be considered approximately constant during some short interval." (p. 1234). This is the reason why, when we use McFarland's model, we set a very short sampleEvery. In addition, the output of the simulation functions contains the simple summary statistic errorMF that can be used to assess the quality of the approximation<sup>16</sup>.

Note that, as the authors point out, approximations are common with stochastic simulation algorithms when there is density dependence, but the advantage of the BNB algorithm compared to, say, most tau-leap methods is that clones of different population sizes are treated uniformly. Mather, Hasty, and Tsimring (2012) further present results from simulations comparing the BNB algorithm with the original direct SSA method and the tau-leaps (see their Fig. 5), which shows that the approximation is very accurate as soon as the interval between samples becomes reasonably short.

# 14.7 Sometimes I get exceptions when running with mutator genes

Yes, sure, the following will cause an exception; this is similar to the example used in 1.3.4 but there is one crucial difference:

```
sd <- 0.1 ## fitness effect of drivers
sm <- 0 ## fitness effect of mutator
nd <- 20 ## number of drivers
nm <- 5 ## number of mutators
mut <- 50 ## mutator effect THIS IS THE DIFFERENCE
fitnessGenesVector <- c(rep(sd, nd), rep(sm, nm))</pre>
```

<sup>&</sup>lt;sup>16</sup>Death rates are affected by density dependence and, thus, it is on the death rates where the approximation that they are constant over a short interval plays a role. Thus, we examine how large the difference between successive death rates is. More precisely, let A and C denote two successive sampling periods, with  $D_A = log(1 + N_A/K)$  and  $D_C = log(1 + N_C/K)$  their death rates. errorMF\_size stores the largest  $abs(D_C - D_A)$  between any two sampling periods ever seen during a simulation. errorMF stores the largest  $abs(D_C - D_A)/D_A$ . Additionally, a simple procedure to use is to run the simulations with different values of sampleEvery, say the default value of 0.025 and values that are 10, 20, and 50 times larger or smaller, and assess their effects on the output of the simulations and the errorMF statistic itself. You can check that using a sampleEvery much smaller than 0.025 rarely makes any difference in errorMF or in the simulation output (though it increases computing time significantly). And, just for the fun of it, you can also check that using huge values for sampleEvery can lead to trouble and will be manifested too in the simulation output with large and unreasonable jumps in total population sizes and sudden extinctions.

Now, simulate using the fitness and mutator specification. We fix the number of drivers to cancer, and we stop when those numbers of drivers are reached. Since we only care about the time it takes to reach cancer, not the actual trajectories, we set keepEvery = NA:

What happened? That you are using five mutator genes, each with an effect of multiplying by 50 the mutation rate. So the genotype with all those five genes mutated will have an increased mutation rate of  $50^5 = 312500000$ . If you set the mutation rate to the default of 1e-6 you have a mutation rate of 312 which makes no sense (and leads to all sorts of numerical issues down the road and an early warning).

Oh, but you want to accumulate mutator effects and have some, or the early ones, have a large effects and the rest progressively smaller effects? You can do that using epistatic effects for mutator effects.

# 14.8 What are good values of sampleEvery?

First, we need to differentiate between the McFarland and the exponential models. If you use the McFarland model, you should read section 14.6 but, briefly, the small default is probably a good choice.

With the exponential model, however, simulations can often be much faster if sampleEvery is large. How large? As large as you can make it. sampleEvery should not be larger than your desired keepEvery, where keepEvery determines the resolution or granularity of your samples (i.e., how often you take a snapshot of the population). If you only care about the final state, then set keepEvery = NA.

The other factors that affects choosing a reasonable **sampleEvery** are mutation rate and population size. If population growth is very fast or mutation rate very large, you need to sample frequently to avoid the "Recoverable exception ti set to DBL\_MIN. Rerunning." issue (see discussion in section 2.4).

## 14.9 What can you do with the simulations?

This is up to you. In section 15.2 we show an example where we infer an Oncogenetic tree from simulated data and in section 1.3 we go over a varied set of scientific questions where OncoSimulR could help.

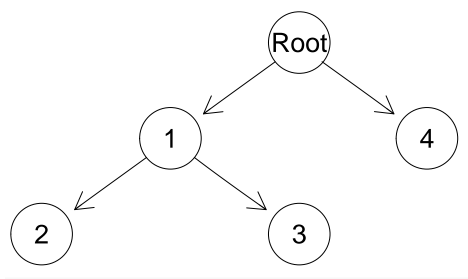
# 15 Using v.1 posets and simulations

It is strongly recommended that you use the new (v.2) procedures for specifying fitness effects. However, the former v.1 procedures are still available, with only very minor changes to function calls. What follows below is the former vignette. You might want to use v.1 because for certain models (e.g., small number of genes, with restrictions as specified by a simple poset) simulations might be faster with v.1 (fitness evaluation is much simpler —we are working on further improving speed).

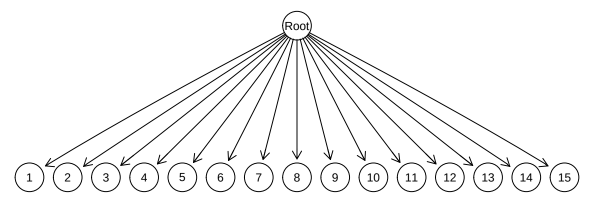
## 15.1 Specifying restrictions: posets

How to specify the restrictions is shown in the help for poset. It is often useful, to make sure you did not make any mistakes, to plot the poset. This is from the examples (we use an "L" after a number so that the numbers are integers, not doubles; we could alternatively have modified storage.mode).

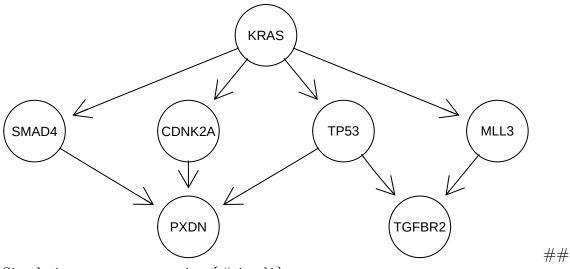
```
## Node 2 and 3 depend on 1, and 4 depends on no one
p1 <- cbind(c(1L, 1L, 0L), c(2L, 3L, 4L))
plotPoset(p1, addroot = TRUE)</pre>
```



```
## A simple way to create a poset where no gene (in a set of 15)
## depends on any other.
p4 <- cbind(OL, 15L)
plotPoset(p4, addroot = TRUE)</pre>
```



Specifying posets is actually straightforward. For instance, we can specify the pancreatic cancer poset in Gerstung et al. (2011) (their figure 2B, left). We specify the poset using numbers, but for nicer plotting we will use names (KRAS is 1, SMAD4 is 2, etc). This example is also in the help for poset:



Simulating cancer progression {#simul1}

We can simulate the progression in a single subject. Using an example very similar to the one in the help:

```
## use poset p1101
data(examplePosets)
p1101 <- examplePosets[["p1101"]]

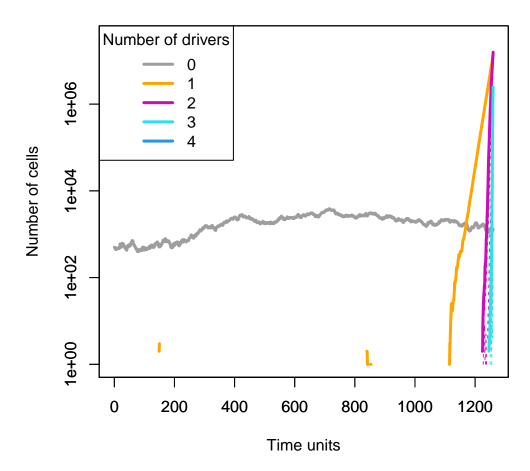
## Bozic Model
b1 <- oncoSimulIndiv(p1101, keepEvery = 15)
summary(b1)
## NumClones TotalPopSize LargestClone MaxNumDrivers</pre>
```

```
## 1 11 29168267 12641599
##
     MaxDriversLast NumDriversLargestPop TotalPresentDrivers
## 1
                                             1
                    4
     \it Final Time \ \it Num Iter \ \it Hitted \it Wall Time \ \it Hitted \it Max Tries \ \it error \it MF
##
## 1
                    3430
                                    FALSE
                                                                  NA
           1371
                                                      FALSE
     minDMratio minBMratio OccurringDrivers
           90909
                        90909 1, 2, 3, 7, 8, 9
## 1
```

The first thing we do is make it simpler (for future examples) to use a set of restrictions. In this case, those encoded in poset p1101. Then, we run the simulations and look at a simple summary and a plot.

If you want to plot the trajectories, it is better to keep more frequent samples, so you can see when clones appear:

```
b2 <- oncoSimulIndiv(p1101, keepEvery = 1)
summary(b2)
     NumClones TotalPopSize LargestClone MaxNumDrivers
##
## 1
                    3.2e+07
                                13713960
            16
##
     MaxDriversLast NumDriversLargestPop TotalPresentDrivers
## 1
                  4
##
     FinalTime NumIter HittedWallTime HittedMaxTries errorMF
## 1
          1260
                  3369
                                FALSE
                                                FALSE
                                                           NA
     minDMratio minBMratio
                                     OccurringDrivers
                     90909 1, 2, 3, 4, 5, 6, 7, 8, 9
## 1
          90909
plot(b2)
```



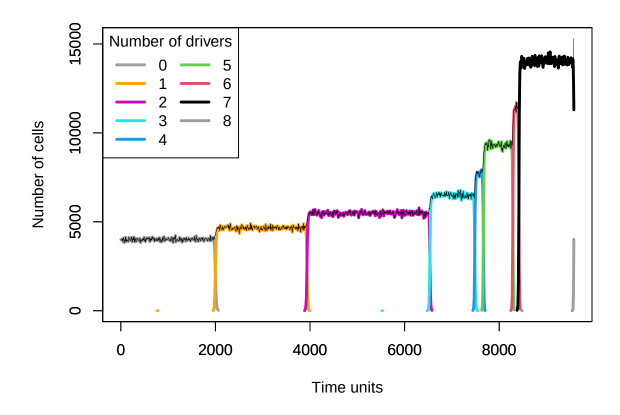
As we have seen before, the stacked plot here is less useful and that is why I do not evaluate that code for this vignette.

```
plot(b2, type = "stacked")
```

The following is an example where we do not care about passengers, but we want to use a different graph, and we want a few more drivers before considering cancer has been reached. And we allow it to run for longer. Note that in the McFL model detectionSize really plays no role. Note also how we pass the poset: it is the same as before, but now we directly access the poset in the list of posets.

(Very rarely the above run will fail to reach cancer. If that happens, execute it again.)

As usual, we will plot using both a line and a stacked plot:





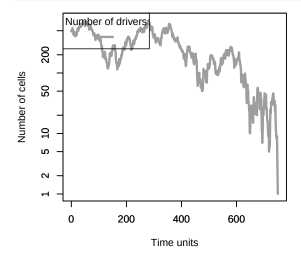
The default is to simulate progression until a simulation reaches cancer (i.e., only simulations that satisfy the detectionDrivers or the detectionSize will be returned).

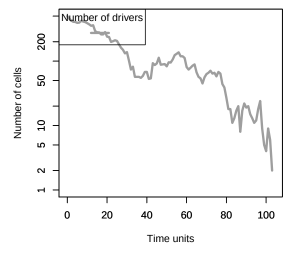
If you use the McFL model with large enough initSize this will often be the case but not if you use very small initSize. Likewise, most of the Bozic runs do not reach cancer. Lets try a few:

```
b3 <- oncoSimulIndiv(p1101, onlyCancer = FALSE)
summary(b3)
##
     NumClones TotalPopSize LargestClone MaxNumDrivers
                            0
## 1
              1
                                          0
##
     {\it MaxDriversLast\ NumDriversLargestPop\ TotalPresentDrivers}
## 1
##
     FinalTime NumIter HittedWallTime HittedMaxTries errorMF
## 1
           751
                    753
                                  FALSE
                                                  FALSE
                                                              NA
     minDMratio minBMratio OccurringDrivers
## 1
          90909
                      90909
b4 <- oncoSimulIndiv(p1101, onlyCancer = FALSE)
summary(b4)
##
     NumClones TotalPopSize LargestClone MaxNumDrivers
## 1
              1
                            0
                                          0
##
     {\it MaxDriversLast} {\it NumDriversLargestPop} {\it TotalPresentDrivers}
## 1
##
     FinalTime NumIter HittedWallTime HittedMaxTries errorMF
                                  FALSE
## 1
            104
                    104
                                                  FALSE
                                                              NA
     minDMratio minBMratio OccurringDrivers
          90909
                     90909
```

Plot those runs:

```
par(mfrow = c(1, 2))
par(cex = 0.8) ## smaller font
plot(b3)
plot(b4)
```





### 15.1.1 Simulating progression in several subjects

To simulate the progression in a bunch of subjects (we will use only four, so as not to fill the vignette with plots) we can do, with the same settings as above:

```
p1 <- oncoSimulPop(4, p1101, mc.cores = 2)
par(mfrow = c(2, 2))
plot(p1, ask = FALSE)
Number of cells
                                                           Number of cells
                                                                  1e+06
      1e+06
            Number of drivers
                                                                        Number of drivers
                         0
                                                                 1e+00
      1e+00
                                                                                  500
            0
                    100
                            200
                                     300
                                             400
                                                                        0
                                                                                            1000
                                                                                                       1500
                          Time units
                                                                                      Time units
Number of cells
                                                           Number of cells
      1e+06
                                                                  1e+06
            Number of drivers
                                                                        Number of drivers
                                                                 1e+00
      1e+00
            0
                    200
                              400
                                       600
                                                 800
                                                                        0
                                                                             200
                                                                                   400
                                                                                          600 800
                                                                                                             1200
```

We can also use stream and stacked plots, though they might not be as useful in this case. For the sake of keeping the vignette small, these are commented out.

Time units

```
par(mfrow = c(2, 2))
plot(p1, type = "stream", ask = FALSE)

par(mfrow = c(2, 2))
plot(p1, type = "stacked", ask = FALSE)
```

### 15.2 Sampling from a set of simulated subjects

Time units

You will often want to do something with the simulated data. For instance, sample the simulated data. Here we will obtain the trajectories for 100 subjects in a scenario without passengers. Then we will sample with the default options and store that as a vector of genotypes (or a matrix of subjects by genes):

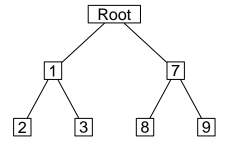
The function samplePop samples that object, and also gives you some information about the output:

```
genotypes <- samplePop(m1)
##
## Subjects by Genes matrix of 100 subjects and 11 genes.</pre>
```

What can you do with it? That is up to you. As an example, let us try to infer an Oncogenetic tree (and plot it) using the *Oncotree* package Szabo and Pappas (2013) after getting a quick look at the marginal frequencies of events:

```
colSums(genotypes)/nrow(genotypes)
## 1 2 3 4 5 6 7 8 9 10 11
## 0.49 0.04 0.02 0.00 0.00 0.00 0.54 0.03 0.04 0.00 0.00

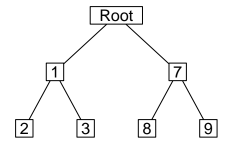
require(Oncotree)
## Loading required package: Oncotree
## Loading required package: boot
ot1 <- oncotree.fit(genotypes)
## The following events had no observed occurances, so they will not be included i
## 4 5 6 10 11
plot(ot1)</pre>
```



Your run will likely differ from mine, but with the defaults (detection size of  $10^8$ ) it is likely that events down the tree will never appear. You can set detectionSize = 1e9 and you will see that events down the tree are now found in the cross-sectional sample.

Alternatively, you can use single cell sampling and that, sometimes, recovers one or a couple more events.

```
genotypesSC <- samplePop(m1, typeSample = "single")</pre>
##
## Subjects by Genes matrix of 100 subjects and 11 genes.
colSums(genotypesSC)/nrow(genotypesSC)
                                          8
           2
                3
                     4
                          5
                               6
                                     7
                                                    10
## 0.59 0.06 0.12 0.00 0.00 0.00 0.65 0.08 0.14 0.00 0.00
ot2 <- oncotree.fit(genotypesSC)</pre>
## The following events had no observed occurances, so they will not be included i
## 4 5 6 10 11
plot(ot2)
```



You can of course rename the columns of the output matrix to something else if you want so the names of the nodes will reflect those potentially more meaningful names.

# 16 Session info and packages used

This is the information about the version of R and packages used:

```
sessionInfo()
## R Under development (unstable) (2020-01-21 r77688)
## Platform: x86_64-pc-linux-gnu (64-bit)
## Running under: Debian GNU/Linux bullseye/sid
##
## Matrix products: default
## BLAS/LAPACK: /usr/lib/x86 64-linux-qnu/libopenblasp-r0.3.7.so
##
## Random number generation:
##
   RNG:
            L'Ecuyer-CMRG
## Normal:
           Inversion
## Sample: Rejection
##
## locale:
   [1] LC CTYPE=en GB.utf8
                                  LC NUMERIC=C
##
   [3] LC TIME=en GB.utf8
                                  LC COLLATE=en GB.utf8
##
    [5] LC_MONETARY=en_GB.utf8
                                  LC_MESSAGES=en_GB.utf8
##
   [7] LC_PAPER=en_GB.utf8
                                  LC_NAME = C
   [9] LC ADDRESS=C
                                  LC TELEPHONE=C
## [11] LC_MEASUREMENT=en_GB.utf8 LC_IDENTIFICATION=C
##
## attached base packages:
## [1] parallel stats
                           graphics grDevices utils
## [6] datasets methods
                           base
##
## other attached packages:
   [1] Oncotree 0.3.3
                            boot_1.3-24
##
                           graph_1.65.0
   [3] igraph_1.2.4.2
##
    [5] BiocGenerics_0.33.0 OncoSimulR_2.17.996
##
    [7] pander_0.6.3
                           bookdown_0.17
##
    [9] BiocStyle_2.15.5
                            rmarkdown_2.1
##
## loaded via a namespace (and not attached):
   [1] gtools_3.8.1
                            tidyselect\_0.2.5
##
                            purrr 0.3.3
##
   [3] x fun \ 0.12
   [5] lattice_0.20-38
##
                            haven_2.2.0
   [7] carData_3.0-3
                            colorspace_1.4-1
   [9] vctrs 0.2.1
##
                            htmltools 0.4.0
## [11] stats4_4.0.0
                            yaml_2.2.0
## [13] rlang_0.4.2
                            pillar_1.4.3
## [15] foreign 0.8-75
                            qlue 1.3.1
## [17] Rgraphviz 2.31.0
                            RColorBrewer 1.1-2
## [19] readxl_1.3.1
                           lifecycle_0.1.0
```

```
## [21] stringr_1.4.0
                            cellranger_1.1.0
## [23] munsell_0.5.0
                            gtable_0.3.0
## [25] zip_2.0.4
                            evaluate_0.14
## [27] knitr 1.27
                            rio 0.5.16
## [29] forcats_0.4.0
                            curl_4.3
## [31] highr_0.8
                            Rcpp_1.0.3
## [33] scales_1.1.0
                            backports_1.1.5
## [35] BiocManager_1.30.10 abind_1.4-5
## [37] farver_2.0.3
                            smatr_3.4-8
## [39] ggplot2_3.2.1
                            hms_0.5.3
## [41] digest_0.6.23
                            stringi_1.4.5
                            dplyr_0.8.3
## [43] openxlsx_4.1.4
## [45] ggrepel_0.8.1
                            grid_4.0.0
## [47] tools_4.0.0
                            magrittr_1.5
## [49] lazyeval_0.2.2
                           tibble_2.1.3
## [51] crayon_1.3.4
                            car_3.0-6
## [53] pkgconfig_2.0.3
                            zeallot_0.1.0
                            data.table_1.12.8
## [55] Matrix_1.2-18
## [57] assertthat 0.2.1
                            R6_2.4.1
## [59] compiler_4.0.0
```

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