Exercise 2: Frequency dependent fitness

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- 2 Functionality
 - Additional functions
 - Modification of the code
- 3 Experiments
 - List of experiments
 - An experiment with bacteria
 - An experiment with cells

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Introduction \rightarrow 2 Frequency dependent fitness

- Game theory.
- OncoSimulR model \rightarrow the fitness of a subpopulation will depend on the relative abundance of the different subpopulations.
- The fitness of each subpopulation is defined as an arbitrary function of the genetic interactions between multiple genes.

Introduction \rightarrow Effects on fitness

- allFitnessEffects function:
 - genoFitness = dataframe
 - First column: genotypes.
 - Second column: expressions for the functions that relate fitness to frequencies of other genotypes.
 - frequencyDependentFitness = TRUE
 - frequencyType = "rel" or frequencyType = "abs"
 - spPopSizes

Introduction \rightarrow Assess fitness

- evalGenotype function:
 - fitnessEffects = allFitnessEffects object
 - genotype
- evalAllGenotypes function:
 - fitnessEffects = allFitnessEffects object

Introduction \rightarrow Perform simulations

• oncoSimulIndiv and oncoSimulPop functions.

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 Box-plot: graphical summary of the distribution of simulations results

compositionPop2()

```
Extract and create a data frame with results from several s
compositionPop2 <- function(objPop, ...) {
  ## Create genotype names
  clon_labels <- c("WT", objPop[[1]]$geneNames)
  ## Extract the information to create a data frame
  listPop \leftarrow vapply(objPop, function(x) tail(x[[1]], 1)[1, -1]
                    as.double(1:length(clon_labels)))
  dfPop <- data.frame("Genotype" = rep(clon_labels,
                                        length(listPop)/length(
                      "N" = c(listPop))
  simul_boxplot2(dfPop, ...)
```

Figure: Code for compositionPop2() function

simul_boxplot2()

```
## Plot box plot (by default same colors as plot.oncosimul type stream)
simul_boxplot2 <- function(df, main = FALSE, xlab = "Genotype", ylab = "N".
                           colors) {
  e \leftarrow ggplot(df, aes(x = Genotype, y = N)) +
    theme(plot.title = element_text(hjust = 0.5, size = 16, face = "bold"),
          axis.title.x = element_text(size = 12, face = "bold"),
          axis.title.y = element_text(size = 12, face = "bold"),
          axis.text.x = element_text(size = 11),
          axis.text.y = element_text(size = 11))
  if (main == FALSE) {
    e + geom_boxplot(aes(fill = Genotype)) +
      stat_summary(fun.y = mean, geom = "point",
                   shape = 18, size = 2.5, color = "#FC4E07") +
      xlab(xlab) + ylab(ylab) +
      scale_fill_manual(values = colors)
      stat_summary(fun.y = mean, geom = "point",
                   shape = 18, size = 2.5, color = "#FC4E07") +
      xlab(xlab) + ylab(ylab) + scale_fill_manual(values = colors)}
  else {
    e + geom_boxplot(aes(fill = Genotype)) +
      stat_summary(fun.y = mean, geom = "point",
                   shape = 18. size = 2.5. color = "#FC4E07") +
      labs(title = main) +
      xlab(xlab) + ylab(ylab) + scale_fill_manual(values = colors)}
```

Figure: Code for simul_boxplot2() function

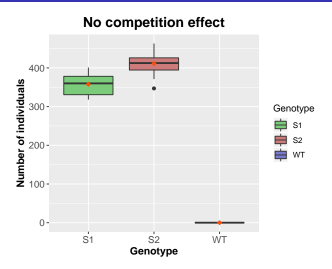


Figure: Box-plot from one of the Lotka-Volterra's example. 20 simulations were made

Stripchart: summary of simulations with oscillating trajectories

```
stripChartPop <- function(dfPop, ylab = "N", ...) {</pre>
  stripchart(dfPop, vertical = TRUE, ylab = ylab, ...)
  f1 <- function(x, num_genotypes) {
    num_genotypes <- length(x)
    i <- 1
    while (num_genotypes > i) {
      segments(x0 = i, x1 = i+1,
               v0 = x[i].
               y1 = x[i+1],
               col = rainbow(5)
      i <- i+1
  apply(dfPop, 1, f1)
meanCompositionPop <- function(objPop, ...) {</pre>
 condi <- c("WT", objPop[[1]]$geneNames)
  ## $pops.by.time contains the times at wich results are taken
  listPop <- lapply(objPop, function(x)
    (colMeans(tail(x$pops.by.time, length(x$pops.by.time[,1])/2)[,-1])))
  dfPop <- data.frame(matrix(unlist(listPop).</pre>
                              ncol = length(condi). byrow = TRUE))
  colnames(dfPop) <- condi
  stripChartPop(dfPop, ...)
  dfPop
```

Figure: stripChartPop() and meanCompositionPop() code

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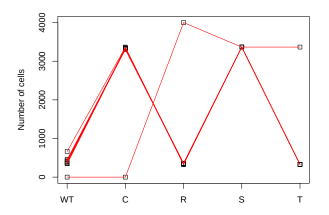
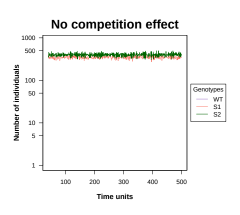


Figure: Example of the graphical summary of an oscillating trajectory

Modification of the code

Legend location



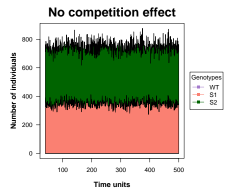


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List of experiments

- Rock-paper-scissors model in bacterial community
- Evolutionary games theory: Hawk and Dove example
- The Lotka-Volterra model of competition between two competing species
- Game Theory with social dilemmas of tumour acidity and vasculature
- Prostate cancer tumour–stroma interactions
- Evolutionary Dynamics of Tumor-Stroma Interactions in Multiple Myeloma

List of experiments

- Rock-paper-scissors model in bacterial community
- ② Evolutionary games theory: Hawk and Dove example
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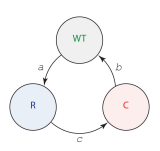
- Title: Local dispersal promotes biodiversity in a real-life game of rock-paper-scissors.
- Authors: Benjamin Kerr, Margaret A. Riley, Marcus W. Feldman, Brendan J. M. Bohannan.
- Three competing species of bacterias with relationships similar to rock-paper-scissors game.

Local dispersal promotes biodiversity in a real game of rock-paper-scissors



- Three types of populations of *E. coli*:
 - Wild-type bacterias (WT): colicin-sensitive bacterias, hence, they are killed by colicin.
 - Colicinogenic bacterias (C): produce colicin toxin, a colicin-specific inmunity protein and a lysis protein which causes partial cell lysis and the release of the colicin.
 - Colicin-resistant bacterias (R): WT bacterias who have mutated getting membrane translocators of colicin.
- Parameters that describe the relationships of WT-C-R community:
 - a: advantage of WT over R ⇒ R have costs in their fitness because the resistance consumes a lot of energy and WT have not this problem.
 - b: advantage of C over WT \Rightarrow C are able to kill WT.
 - c: advantage of R over C \Rightarrow R are resistant to colicin produced by C.

In summary, we have the relationships of rock-paper-scissors game:



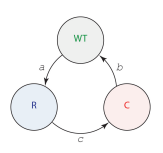
And the resulting equations:

$$W(WT) = 1 + af_3 - bf_2$$

 $W(C) = 1 + bf_1 - cf_3$
 $W(R) = 1 + cf_2 - af_1$

where f_1 , f_2 and f_3 are the frequencies of WT, C and R, respectively.

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And the resulting equations:

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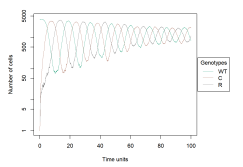
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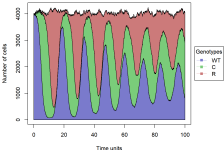
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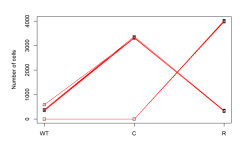
allFitnessEffect and oncoSimulIndiv functions

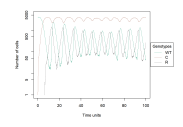
Case 1: a = b = c = 1

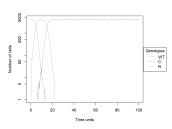




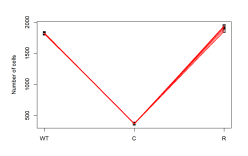
Case 2:
$$a = 10$$
, $b = c = 1$

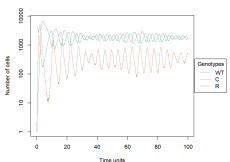






Case 3:
$$a = 1$$
, $b = c = 5$





Tumor-Stroma Interactions

- **Title:** Evolutionary Dynamics of Tumor-Stroma Interactions in Multiple Myeloma.
- Authors: Javad Salimi Sartakhti, Mohammad Hossein Manshaei, Soroosh Bateni, Marco Archetti.
- Cancer cells and stromal cells cooperate by exchanging diffusible factors.
 - Frequency-dependent selection that can be studied in the framework of evolutionary game theory.

Tumour-Stroma Interactions: payoff functions

- 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.
- The payoff for strategy P_j is:

$$\pi_{P_j}(M_1,\ldots,M_n)=\frac{(M_j+1)\times c_j}{M}r_{j,j}+\sum_{i=1,i\neq j}^n\frac{M_i\times c_i}{M}r_{j,i}-c_j.$$



Tumour-Stroma Interactions: dynamics

- Malignant plasma cells.
- Osteoblasts.
- Osteoclasts.
- Growth factors:
 - Autocrine effects.
 - Paracrine effects.

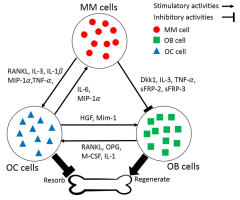
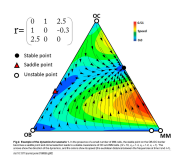


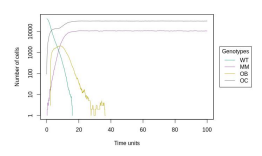
Fig 1. Bone remodeling in multiple myeloma. Multiple myeloma cells (MM) produce growth factors that activate osteoclasts (OC), which increase bone rescoption, or that inhibit osteoclast (OB) differentiation. OC and OB secrete growth factors that affect each other and MM cells.

doi:10.1371/journal.pone.0168856.g001

Tumour-Stroma Interactions: Scenario 1

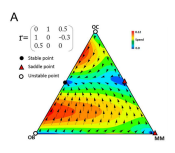
- $c_1 < c_2 < c_3$ (a common occurrence in multiple myeloma).
- In the presence of a small number of MM cells, the stable point on the OB-OC border becomes a saddle point and clonal selection leads to a stable coexistence of OC and MM cells.

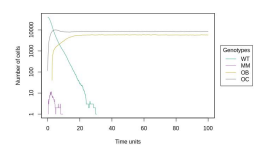




Tumour-Stroma Interactions: Scenario 2

- $c_1 = c_2 = c_3$.
- The game has one polymorphic stable point between OB and OC. In this case, clonal selection leads to the regular OC-OB balance and prevents invasion of MM cells.





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