# Gillespie Algorithm

#### Initialisation

Firstly, we need to install required packages.

```
require(GillespieSSA)
require(ggplot2)
require(dplyr)
require(foreach)
require(reshape2)
```

To do this you should use function install.packages(package\_name).

# Gene expression model.

We will consider simple GEM defined as:

```
dm/dt = Sm - km * m dn/dt = Sn * m - kn * n
```

To simulate model using Gillespie Algorithm we will use ssa function implemented in package GillespieSSA. Let's read the ssa's documentation ?ssa.

To simulate model we need to define:

- nu state change matrix (stoichometric matrix)
  - rows corresponds to variables
  - columns corresponds to reactions
  - each element e\_{i,j} of matrix tells what is an effect of reaction j on variable i
- a propensity vector prawdopodobienstwo przejscia pomiedzy stanami reakcji

```
nu <-
    matrix(
        c(1, -1, 0, 0,
            0, 0, 1, -1),
    nrow = 2,
    byrow = TRUE)
a <-
    c("Sm",
        "km*m",
        "sn*m",
        "kn*n")</pre>
```

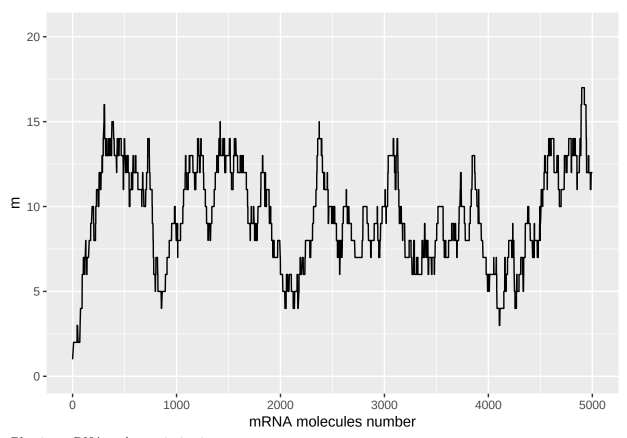
Moreover we need to initiate parameters and variables state:

- x0 initial variables state
- param vector of parameters values
- tf time of simulations

```
x0 \leftarrow c(m = 1, n = 0)
parms \leftarrow c(Sm = 0.05, km = 0.005, Sn = 0.05, kn = 0.001)
tf \leftarrow 5000
```

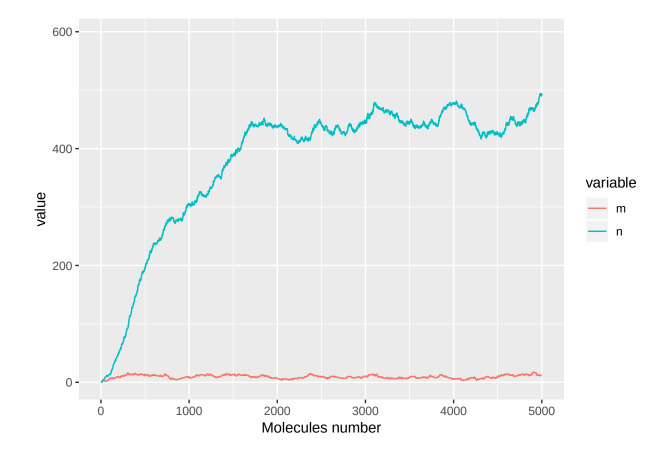
Model simulation:

```
method <- "D"
simName <- "Gene Expression Model"</pre>
out \leftarrow ssa(x0 = x0,
            a = a,
             nu = nu,
             parms = parms,
             tf = tf,
             method = method,
             simName = simName,
             verbose = FALSE,
           consoleInterval = 1)
data <- data.frame(out$data)</pre>
colnames(data)[1] <- "t"</pre>
head(data)
##
                          t m n
## timeSeries 0.000000 1 0
## X 9.795561 2 0
## X.1 13.575579 2 1
## X.2 13.860197 2 2
## X.3 16.258750 2 3
## X.4 20.150344 2 4
Plotting mRNA
g <-
  ggplot(
    data = data,
    mapping = aes(x = t, y = m)) +
  geom_line() +
  xlab("mRNA molecules number") +
  coord_cartesian(ylim = c(0, 1.2*max(data$m)))
print(g)
```



#### Plotting mRNA and protein in time

```
data <- data.frame(out$data)
colnames(data)[1] <- "t"
data.long <-
    data %>%
    reshape2::melt(id = "t")
g <-
    ggplot(
    data = data.long,
    mapping = aes(x = t, y = value, group = variable, color = variable)) +
    geom_line() +
    xlab("Molecules number") +
    coord_cartesian(ylim = c(0, 1.2*max(data$m, data$n)))
print(g)</pre>
```



## Task 1

Consider GEM with different initiate state values.

- Initiate in steady state
- Initiate state m = 1, n = 0
- Initiate state m = 10, n = 0

Prepare code that simulates model multiple times and plot trajectories of molecules numbers.

- Calculate and plot mean and standard deviation of molecules in time. For mRNA compare results in steady state with analytical solution of master equations.
- Calculate time of convergence to the steady state.
- Plot mRNA vs protein

## Task 2

Lottka-Volter Model

- R1:  $A + X \rightarrow 2X$
- R2: X + Y -> 2Y
- R3: Y -> \*

 $X-rabbits\ Y-foxes$ 

'reaction' rates: k1, k2, k3

A - paramter

- Calulcate the propensity function and stechiometric matrix
- Caluclate steady state
- Simulate model multiple times starting from different states. What is a time of convergance ?
- Prepare plot presenting population of rabbits vs foxes

### Task 3

Zrób symulację następującego układu:

 $A + B \rightarrow C$ 

 $\mathrm{D} \to \mathrm{A} + \mathrm{B}$ 

 $\to E$ 

 $E\,+\,C\,\to\,D$ 

 $A \rightarrow *$ 

 $\mathrm{B} o *$ 

 $C \rightarrow *$ 

 $D \rightarrow *$ 

 $E \rightarrow *$ 

i znajdź taki rozkład prawdopodobieństw poszczególnych reakcji, który skutkuje dwoma scenariuszami: układ znajduje się mniej więcej w stanie stacjonarnym układ rośnie w nieskończoność Pokaż wykresy stężeń dla wszystkich związków.

Jako stan początkowy przyjmij losowe ilości cząsteczek wszystkich związków z przedziału <2,10>.