EMERGENT THRESHOLDS IN GENETIC REGULATORY NETWORKS: PROTEIN PATTERNING IN DROSOPHILA MORPHOGENESIS

RUI DILÃO AND DANIELE MURARO

Nonlinear Dynamics Group, Instituto Superior Técnico,
Av. Rovisco Pais, 1049-001 Lisbon, Portugal.

rui@sd.ist.utl.pt, muraro@sd.ist.utl.pt

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ABSTRACT. We present a general methodology in order to build mathematical models of genetic regulatory networks. This approach is based on the mass action law and on the Jacob and Monod operon model. The mathematical models are built symbolically by the Mathematica software package GeneticNetworks. This package accepts as input the interaction graphs of the transcriptional activators and repressors and, as output, gives the mathematical model in the form of a system of ordinary differential equations. All the relevant biological parameters are chosen automatically by the software. Within this framework, we show that threshold effects in biology emerge from the catalytic properties of genes and its associated conservation laws. We apply this methodology to the segment patterning in Drosophila early development and we calibrate and validate the genetic transcriptional network responsible for the patterning of the gap proteins Hunchback and Knirps, along the antero-posterior axis of the Drosophila embryo. This shows that patterning at the gap genes stage is a consequence of the relations between the transcriptional regulators and their initial conditions along the embryo.

1. Introduction

A genetic regulatory network is an ensemble of interactions in a biological process involving proteins, genes and mRNAs. The interactions between different proteins and genes can be done by transcriptional activation and repression at the level of the genes, by protein-protein interactions, and by protein-mRNA interactions.

A genetic regulatory networks is described by a graph where vertices represent genes, proteins, enzymes or other chemical substances. The edges represent transformations, e. g., phosphorylation and dephosphorylation, or activation and inhibitory actions through transcription regulators.
More precisely, a genetic regulatory networks is described by a double graph $G = (V, E_a, E_r)$, where $V$ is the set of vertices or nodes of the graph and $E_a$ and $E_r$ are two sets of ordered pairs of vertices of the double graph. Each ordered pair of vertices defines the activation or the repression mechanism of the first node over the second. In classical graph theory, $G = (V, E_a)$ and $G = (V, E_r)$ are two graphs with a common set of vertices. For example, in Figure 1, we show the graph of a genetic network associated with the production of the proteins Bicoid (BCD), Hunchback (HB), Knirps (KNI) and Tailless (TLL) in *Drosophila* early development, (Alves and Dilão, 2006). In this example, we have $V = \{bcd, hb, BCD, HB, KNI, TLL\}$, $E_a = \{(bcd, BCD), (hb, HB), (BCD, HB), (BCD, KNI)\}$ and $E_r = \{(HB, KNI), (KNI, HB), (TLL, KNI)\}$.

![Figure 1. Graph describing the genetic network associated with the production of proteins Bicoid (BCD), Hunchback (HB), Knirps (KNI) and Tailless (TLL) in *Drosophila* early development. mRNAs *hunchback* and *bicoid* are represented by hb and bcd respectively. Arrows represent activations and are listed in the set of ordered pairs $E_a$. Lines with perpendicular endings represent repressions and are listed in the set $E_r$.](image)

The graph of Figure 1 has a clear biological meaning. It expresses the fact that BCD is a transcriptional activator of both HB and KNI, HB and KNI proteins both repress each other, and TLL is a repressor of KNI. The vertices of the graph of Figure 1 can represent mRNAs, as in the case of hb and bcd, or proteins, as in
the case of BCD and TLL, or genes and proteins simultaneously, as in the case of 
HB and KNI.

Here we propose a set of rules in order to construct the model equations 
associated with a genetic regulatory networks described by a double graph $G = 
(V, E_a, E_b)$. This paper is an attempt to delineate a methodological approach for 
the construction of mathematical models of genetic regulation from the principles 
of chemical kinetics and chemical bound. In the literature, it is often found ex-
amples of mathematical models of biological systems described by different sets 
of equations and characterized by different sets of parameters that are difficult to 
interpret and measure. Making qualitative predictions with these different models 
has a limited predictable value. For a review on the different approaches see, for 
example, (Klipp et al. 2005) and Jong (2002).

From the point of view of predictability, it is important that mathematical model 
could be derived from first principles, leading to a meaningful identification of the 
biological parameters, their calibration and validation with experimental data.

In the construction of models for generic regulatory networks, we assume that 
models can be built with rate equations reflecting a mean field view of the stochas-
tic random motion occurring at the molecular scale. This mean field approach, 
also called mass action law, is derived from the probabilistic collision laws occur-
ring at the molecular scale. The models originated from this view are described by 
ordinary differential equations with polynomial vector fields, (van Kampen 1992).

One of the advantages of the mass action law approach is that the mean field 
rate equations have a direct microscopic interpretation, being associated with the 
collision mechanism that are in the origin of every reactive process. For model re-
finement, fluctuations can also be studied through the corresponding master equa-
tion. From the experimental point of view, microbiology techniques are strongly
anchored in the mass action law or mean field approach. For example, genome sequencing is only possible in the framework of the mean field approach.

For genetic regulatory networks described by graphs with a large number of vertices, and a complex structure of edges, the rate equations describing the evolution in time of concentrations are in general difficult to build, and are critically dependent on the assumptions done about the biological and chemical interactions involved. During the development of these complex models, it is often necessary to test different configurations changing the parameters and the initial conditions. Writing by hand all of these information is both time-consuming and error-prone.

In order to perform these tasks automatically, we have developed two Mathematica software packages, Kinetics and GeneticNetworks, that execute the symbolic computations associated with the construction of the model equations for a genetic regulatory network. The result of the analysis is in symbolic form, and can be used in Mathematica or in C for further processing and numerical integration. The software packages GeneticNetworks and Kinetics are freely distributed from the site “https://sd.ist.utl.pt”.

The Kinetics package implements the mass action law in its polynomial exact form, computing symbolically the associated rate equations and conservation laws. The package GeneticNetworks implements a particular model for protein-gene regulation. This model for the protein-gene regulation is based on the operon model of Jacob and Monod [Jacob and Monod 1961], and its basic properties have been previously introduced in Alves and Dilão [2005].

This paper is organized as follows. In section 2 we briefly review the mass action law of chemical kinetics and we introduce the collision graphs associated with the mass action law. We derive the basic mass action rate equations. A special emphasis is done on mass action conservation laws, an important feature
that is in the very foundations of threshold effects in biology. In other approaches, threshold do not result as emergent phenomena, but must be imposed through “ad-hoc” regulatory functions (see for example (Klipp et al. 2005, pp. 237) or (Jong 2002)).

In section 3, we describe the mechanism of genetic regulation based on the Jacob and Monod operon model, (Jacob and Monod 1961), and we introduce the modelling assumptions for the construction of the mathematical models of genetic networks described by double graphs.

In section 4, we give an overview of the GeneticNetworks software package. In section 5, we show three different applications of genetic regulatory networks. In the first application, we show, with a very simple example of autoregulation, that the conservation law constant is a bifurcation parameter for the regulation model, inducing a threshold effect in the production of proteins. In the second example, we give a genetic regulatory network inducing a localized spiky pattern. In the third example, we analyze the experimental data associated with the KNI and HB inhibitory cross regulation in Drosophila early development, described by the double graph of Figure 1, and we calibrate this model with the experimental data.

In the final section, we summarize the main biological conclusions of the paper.

2. The mass action law framework of chemical kinetics

In general, an ensemble of chemical reactions is represented by the following collision diagram,

\[ \nu_{i1}A_1 + \cdots + \nu_{im}A_m \xrightarrow{r_i} \mu_{i1}A_1 + \cdots + \mu_{im}A_m \]

where \( i = 1, \ldots, n \). The \( A_j \), for \( j = 1, \ldots, m \), represent chemical substances, as for example, \( A_j = \text{H}_2\text{O} \). The constants \( \nu_{ij} \) and \( \mu_{ij} \) are the stoichiometric coefficients,
in general, non-negative integers, and the constants $r_i$ are the rate constants. If $\nu_{ij} = \mu_{ij} > 0$, the corresponding substance $A_j$ is a catalyst and, if $\mu_{ij} > \nu_{ij} > 0$, $A_j$ is an autocatalyst. In the diagram (1), there are $m$ chemical substances and $n$ rate constants or chemical reactions.

Under the hypothesis of homogeneity of the solution where reactions occur, the mass action law asserts that the time evolution of the concentrations of the chemical substances is described by the system of ordinary differential equations,

\begin{equation}
\frac{dA_j}{dt} = \sum_{i=1}^{n} r_i (\mu_{ij} - \nu_{ij}) A_{i}^{\nu_{i1}} \cdots A_{m}^{\nu_{im}}
\end{equation}

where $j = 1, \ldots, m$, and we use the same symbol to represent both the chemical substance and its concentration. The rate equations (2) are derived under the following assumptions: (i) chemical reactions, when they occur, are due to elastic collisions between the reactants, (ii) homogeneity of the reacting substances in the solution, and (iii) thermal equilibrium of the solution. All the kinetics aspects related with the dependence of the velocity of the reactions on the temperature or pressure are contained in the rate constants $r_i$. For details see van Kampen (1992).

At the atomic and molecular scale, chemical reactions between molecules can occur only if molecules collide or approach each other to small distances where bounding forces become meaningful. These chemical bounding forces are of electrical or quantum origin, and at distances larger than the mean free path they become less important when compared with the kinetics associated with the molecular motion. As chemical reactions only occur if the chemical substances involved collide, the vector fields associated with the right hand side of (2) are in general quadratic, representing binary collisions. Higher order polynomial vector fields are
possible but, at the microscopic level, they are associated to triple or higher order collisions, a situation that occurs with a very low probability. Therefore, we will restrict our examples to models with two-body collisions.

The equations (2) can also be written in the matrix form,

\[
\frac{dA}{dt} = \Gamma \omega(A)
\]

where \(\Gamma\) is a \(n \times m\) matrix, \(A^T = (A_1, \ldots, A_m)\), and,

\[
\omega(A) = \begin{pmatrix}
 r_1(\mu_1 j - \nu_1 j)A_{\nu_11}^1 \cdots A_{\nu_1m}^m \\
 \vdots \\
 r_n(\mu_n j - \nu_n j)A_{\nu_n1}^1 \cdots A_{\nu_nm}^m
\end{pmatrix}
\]

In general, \(n \neq m\), and the equations in system (2) are not all independent. Let us denote by \(r\) the rank of the matrix \(\Gamma\). The dimension of the null space of \(\Gamma\) relates with its rank by, \(\dim(\text{Null}(\Gamma)) + r = m\) (number of columns of \(\Gamma\)). Let \(v_1, \ldots, v_{m-r}\) be a basis of the Null space of \(\Gamma\), then, \(\Gamma v_k = 0\), for \(k = 1, \ldots m-r\). So, by (3), we have,

\[
\frac{dA}{dt}.v_k = \frac{d}{dt}(A.v_k) = (\Gamma \omega).v_k = \omega.(\Gamma v_k) = 0
\]

Hence, associated to the differential equations (2), we have the conservation laws,

\[
A.v_k = \text{cons}
\]

where, \(k = 1, \ldots m-r\).

The Mathematica software package Kinetics calculates the rate equations (2) describing the time evolution of the concentrations of the substances involved in
the reactions described by the collision diagram (1). The package calculates also
the corresponding conservations laws (5).

The input of the package is the ensemble of chemical reactions, and the output
of the package is the set of differential equations derived by the mass action law.
The output can be later analyzed and studied by the analytical and numerical
tools provided by Mathematica. In order to avoid long development times, the
names of the rate constants are chosen automatically by the program.

The package Kinetics has the usual help commands, and we provide the Mathematica
notebook KineticsTest.nb with several self-explanatory examples and com-
putations.

For example, let us now describe a simple protein production model with Kinetics. The molecular biology dogma asserts that genes are the templates for
protein production, and the standard mechanism for protein production can be
represented by the collision diagrams,

\[
\begin{align*}
\text{Gene} + \text{Polymerase} & \xrightarrow{r_1} \text{Gene} + \text{Polymerase} + mRNA \\
mRNA & \xrightarrow{r_2} mRNA + \text{Protein} \\
mRNA & \xrightarrow{r_3} \\
\text{Protein} & \xrightarrow{r_4} 
\end{align*}
\]

(6)

Using the symbols $G$, $Pol$, $R$ and $P$ to represent gene, polymerase, mRNA and
protein concentrations, respectively, the collision equations (6) are the input for
Kinetics, with the syntax,

\[
\text{input}=\{G + Pol \to G + Pol + mRNA, mRNA \to mRNA + P, mRNA \to W1, P \to W2\}
\]

where $W1$ and $W2$ are waist products.
For the collision mechanism (6), the rate equations for the protein concentration, and calculated by the package *Kinetics*, are,

\begin{align*}
G' &= 0 \\
Pol' &= 0 \\
R' &= r_1 G.Pol - r_3 R \\
P' &= r_2 R - r_4 P
\end{align*}  

(7)

and the rate constants have been chosen automatically by the software package. In this model, genes and mRNAs are catalysts, and these equations have the exact solutions,

\begin{align*}
G(t) &= G(0) \\
Pol(t) &= Pol(0) \\
R(t) &= R(0) e^{-r_3 t} + \frac{r_1 G(0) Pol(0)}{r_3} (1 - e^{-r_3 t}) \\
P(t) &= G(0) Pol(0) \frac{r_1 r_2}{r_3 r_4} \\
&+ \left( P(0) + \frac{R(0) r_2 r_4 - r_1 r_2 G(0) Pol(0)}{r_4 (r_3 - r_4)} \right) e^{-r_4 t} \\
&+ \frac{R(0) r_3 - r_2 G(0) Pol(0)}{r_3 (r_4 - r_3)} e^{-r_3 t}
\end{align*}  

(8)

To simplify the model equations of protein production and maintaining the catalyst properties of genes, in the following, instead of the collision mechanism (6), we use the simplified or reduced kinetic mechanism,

\begin{align*}
\text{Gene} \xrightarrow{s_1} \text{Gene} + \text{Protein} \\
\text{Protein} \xrightarrow{s_2}
\end{align*}  

(9)
To the reactions (9) correspond the rate equations,

\[ G' = 0 \]
\[ P' = s_1 G - s_2 P \]

The rate equations (10) have the solutions,

\[ G(t) = G(0) \]
\[ P(t) = G(0) \frac{s_1}{s_2} + (P(0) - G(0) \frac{s_1}{s_2}) e^{-s_2 t} \]

Comparing the protein solutions in (11) and (8), we conclude that the steady state of protein of both models is unique and is proportional to the gene concentration. The proportionality constant is different for both models, depending on the rates of the reactions involved. The steady state \( G(0) \frac{s_1}{s_2} \) of model (9) has a direct biological meaning: \( s_1 \) is the rate of protein production and \( s_2 \) is the rate of protein degradation, and \( G(0) \) is the initial gene concentration.

In the following, and in order to simplify the description of the transcriptional regulation of proteins, we will adopt the mechanism (9) in order to describe protein production.

In both rate equations (10) and (7), the concentration of gene is constant along time, and therefore the gene concentration is a conservation law. In the following, we will show that the linear conservation laws of the form (5), will have an important role in the determination of steady states and in bifurcations associated with threshold effects.
3. A mass action framework to describe genetic regulatory networks based on the protein-gene interaction

In the previous section, we have described a basic model model for the production of proteins, in the framework of the mass action law. Based on this framework, we generalize this view in order to include the case of transcriptional regulation by proteins.

In order to keep general the approach presented here and to maintain the biological reality of model parameters, we make the following basic modeling assumptions:

1) In order to describe quantitatively the protein production (concentration) within the molecular biology dogma, we only consider genes and proteins. Intermediate substances in the regulatory mechanism like catalysts and mRNAs are not considered. A model of protein production has been presented and analyzed at the end of the previous section.

2) The regulation of protein production by the template gene is based on the Jacob and Monod operon model [Jacob and Monod, 1961]. Namely, every gene has associated a certain number of binding sites where transcription factors can bind — activators or repressors, Figure 2. The regulation of activations and repressions occurs only through the binding sites. For a given double graph of interactions, the number of binding sites of a gene is determined by the number of edges that end up in the corresponding graph node.

3) Transcription factors are the proteins associated with the vertices that activate or inhibit the production of other proteins. The vertex of a graph represent a transcription factor only if it is the initial point of a edge of activation or inhibition. If a vertex has incoming and outcoming edges of any type, this vertex represents a protein and a gene with several binding states.
4) Each transcription factor has its own binding site in the gene strand, or each gene has only one binding site for all the regulators. Both cases are treated separately in the model. We assume that when at least one activator is bound to a gene, the transcription is activated with a particular production rate for each combinatorial possibility of all the remaining binding sites.

For example, in the double graph of the biological mechanism of Figure 1, we have the following chemical substances,

\begin{align*}
\text{bcd, hb} & \quad \text{mRNAs} \\
\text{BCD, HB, TLL} & \quad \text{proteins} \\
\text{HB}_{0}^{0}, \text{HB}_{0}^{BCD}, \text{HB}_{KNI}^{0}, \text{HB}_{KNI}^{BCD} & \quad \text{operons} \\
\text{KNI}_{0,0}^{0}, \text{KNI}_{0,0}^{BCD}, \text{KNI}_{HB,0}^{0}, \text{KNI}_{HB,0}^{BCD} & \quad \text{operons} \\
\text{KNI}_{0,TLL}^{0}, \text{KNI}_{0,TLL}^{BCD}, \text{KNI}_{HB,TLL}^{0}, \text{KNI}_{HB,TLL}^{BCD} & \quad \text{operons}
\end{align*}

The description of the time evolution of protein concentrations of the mechanisms of Figure 1 involves one rate equation for each substance in (12), except eventually for bcd and hb. As proteins are produced from a gene template, the symbol associated to each vertex of the graph represents a protein. The operon states are represented by the same symbol with superscripts and lowerscripts. The superscripts positions indicate the binding or unbinding of transcriptional activators. The lowerscripts positions indicate the binding or unbinding of transcriptional repressors.

The model associated with a given double graph contains the rate equations for the proteins and the operons in its different states. We also assume by default that proteins always degrade and genes are autocatalytic substances that never degrade. The first assumption implies that protein concentrations remain bounded in time,
Figure 2. Jacob and Monod operon model for the regulation of protein production. The transcription is regulated by the activators and the repressors binding to the binding sites of the gene. Figure adapted from [Alves and Dilão, 2005].

and the second assumption implies the existence of a conservation law for the concentration of the operon states.

4. The GeneticNetworks software package

GeneticNetworks is a software package that generates the rate equations for the concentrations of genes and proteins in a regulatory network. The starting point is the double graph of activations and repressions, $G = (V, E_a, E_r)$. The inputs for GeneticNetworks are two strings, activation and repression, that describe the transcriptional activations and repressions of proteins on genes. In the graph, the same symbol is used to denote both a gene and the corresponding produced protein. As we have seen in (12), the set of variables for the regulation model is constructed with the vertex symbols and the associated operon states.

For example, using as input for GeneticNetworks the interaction strings,

\[
\begin{align*}
\text{activation} &= \{A \rightarrow B\} \\
\text{repression} &= \{R \rightarrow B\}
\end{align*}
\]

the double graph of the genetic network (13) is shown in Figure 3. In this case, the double graph $G = (V, E_a, E_r)$ is characterized by the sets, $V = \{A, B, R\}$, $E_a = \{(A, B)\}$ and $E_r = \{(R, B)\}$.
Figure 3. Double graph associated with the input strings (13) for the GeneticNetworks software package.

In the interaction mechanism (13), protein A activates gene B, and protein R represses gene B. Therefore, the variables of the mechanism (13) are,

\[ A, B, R \] proteins
\[ B_0^B, B_0^A, B_R^A, B_R^R \] operons

The following functions are defined in the GeneticNetworks package:

- NetworkGraph, ManipulateGraph
- Reactions, ReactionsOneSite, ReactionGraph
- SubstanceNames, SubstanceVariables, SubstanceInitialConditions
- ParameterNames, ParameterInput
- Equations
- ConservationLaws

With these functions, we calculate the model equations associated with the input strings (13), calculate automatically the number of variables of the model, define all the relevant parameters and calculate the rate equations. For example, to the genetic regulatory network (13), the GeneticNetworks package associate the
collision diagrams,

\[ A + B_0^0 \xrightleftharpoons[b_a]{b_{-a}} B_A^0 \]

\[ R + B_0^0 \xrightleftharpoons[b_{r}]{b_{-r}} B_R^0 \]

\[ R + B_0^A \xrightleftharpoons[b_{-r}]{b_{r}} B_R^A \]

\[ A + B_R^0 \xrightleftharpoons[b_{-a}]{b_{a}} B_R^A \]

\[ B_0^A \xrightarrow{(p^A_A)_B} B_0^A + B \]

\[ B_R^A \xrightarrow{(p^A_A)_B} B_R^A + B \]

\[ B \xrightarrow{d_B} \]

To these collision diagrams, we have the mass action law rate equations,

\[ A' = -b_a A . B_R^0 + b_{-a} B_R^A - b_a A . B_0^0 + b_{-a} B_0^A \]

\[ B' = (p_A^A)_B B_R^A + (p_0^A)_B B_0^A - d_B B \]

\[ R' = -b_r R . B_0^A + b_{-r} B_R^A - b_r R . B_0^0 + b_{-r} B_R^0 \]

\[ (B_0^0)' = -b_a A . B_0^0 + b_{-a} B_0^A - b_r R . B_0^0 + b_{-r} B_R^0 \]

\[ (B_A^0)' = b_a A . B_0^0 - b_{-a} B_0^A - b_r R . B_A^0 + b_{-r} B_R^A \]

\[ (B_R^0)' = -b_a A . B_R^0 + b_{-a} B_R^A + b_r R . B_0^0 - b_{-r} B_R^0 \]

\[ (B_A^R)' = b_a A . B_R^0 - b_{-a} B_R^A + b_r R . B_0^A - b_{-r} B_R^A \]

and the conservation law,

\[ (B_0^0(t) + B_A^0(t) + B_R^0(t) + B_A^R(t) = B_0^0(0) + B_A^0(0) + B_R^0(0) + B_A^R(0) \]

From the conservation law (17), we can eliminate one of the equations in (16). In this genetic network, we have assumed that the protein concentrations of \( A \) and \( R \) are constant along time.
The rate equations (16) define a mass action law based model for the genetic regulatory network of Figure 3.

In the implementation of GeneticNetworks, we have two possible modeling choices. In one choice, each different regulator has its own binding site, and the model diagrams (15) have been constructed with this assumption. For the second choice, we consider that there is only one binding site in the operon where all the regulators bind. In this case, the collision diagrams associated with the genetic network (13) and calculated in the GeneticNetworks package are,

\[
\begin{align*}
A + B_0 &\xrightleftharpoons{b_a}{b_{-a}} B_A \\
R + B_0 &\xrightleftharpoons{b_r}{b_{-r}} B_R \\
B_A &\xrightarrow{p_B} B_A + B \\
B &\xrightarrow{d_B} \\
\end{align*}
\]  

(18)

By the mass action law, the ReactionsOneSite command leads to the rate equations,

\[
\begin{align*}
A' &= b_{-a}B_A - b_a A.B_0 \\
B' &= p_B B_A - d_B B \\
R' &= b_{-r}B_R - b_r B_0.R \\
(B_0)' &= -b_a A.B_0 + b_{-a} B_A - b_r B_0.R + b_{-r} B_R \\
(B_A)' &= b_a A.B_0 - b_{-a} B_A \\
(B_R)' &= b_r B_0.R - b_{-r} B_R \\
\end{align*}
\]  

(19)

The equations (18) have the conservation law,

\[
B_0(t) + B_A(t) + B_R(t) = B_0(0) + B_A(0) + B_R(0)
\]  

(20)
The two models (15) and (18) for the genetic network (13) are different and these two choices are implemented in the GeneticNetworks package. For the dynamical analysis of a particular case of the distinction between the two models (15) and (18), see (Alves and Dilão, 2005). Below, we will show with a specific example that these two different regulation choices lead to qualitatively and quantitatively similar results.

5. Applications

5.1. An emerging threshold in the dynamics of a self-activating protein.
As an application of the rules describing a genetic regulatory network just introduced, we discuss now the basic role of the conservation laws in the occurrence of threshold effects in regulation mechanisms. We study the case of a self-activating protein, where the produced protein activates its own production, Figure 4.

![Regulation graph describing a self-activating protein.](image)

**Figure 4.** Regulation graph describing a self-activating protein.

The simplest self-activating genetic network is described by the input tables,

\[
\text{activation} = \{A \rightarrow A\} \\
\text{repression} = \{\}.
\]
The reactions and the parameters involved in this activation can be obtained by the
*GeneticNetworks* command `Reactions`, followed by the command `ReactionGraph`,

\[
\begin{align*}
A^0 + A & \xrightleftharpoons{a_a}{a_{-a}} A^A \\
A^A & \xrightarrow{p_A} A^A + A \\
A & \xrightarrow{d_A} 
\end{align*}
\]  

(21)

where \(A^0\) and \(A^A\) are operon states and \(A\) is the corresponding protein.

With the command `Equations`, we get the rate equations,

\[
\begin{align*}
A' &= a_a A^0 A^A + a_{-a} A^A + p_A A^A - d_A A \\
(A^0)' &= -a_a A^0 A^A - a_{-a} A^A \\
(A^A)' &= a_a A^0 - a_{-a} A^A
\end{align*}
\]

(22)

Finally, with the command `ConservationLaws`, we find,

\[
A^0(t) + A^A(t) = A^0(0) + A^A(0) = c
\]

(23)

where \(c\) is a constant.

Introducing (23) into (22), the independent set of rate equations describing the
process (21) is,

\[
\begin{align*}
A' &= a_a A^0 A^A + a_{-a}(c - A^0) + p_A(c - A^0) - d_A A \\
(A^0)' &= -a_a A^0 A^A + a_{-a}(c - A^0)
\end{align*}
\]

(24)

We analyze now the steady state and the phase space structure of the solutions
of equations (24). Equations (24) have two steady states with coordinates,

\[(A^0_{(1)}, A_{(1)}) = (c, 0)\]
and,

\[(A_0(2), A(2)) = \left( \frac{d_A a_{-a}}{p_A a_a}, \frac{cp_a a_a - d_A a_{-a}}{d_A a_a} \right)\]

As the coordinate of the two fixed points are dependent of \(c\), by (23), the steady state coordinates are dependent of the initial concentrations of the operon.

Let \(J(i)\), \(i = 1, 2\) be the Jacobian of equation (24) evaluated at the fixed points. As,

\[\det J(1) = -\det J(2)\]

and,

\[\det J(1) < 0 \iff a_{-a} d_A < ca_a p_A\]

then, we have,

\[
\begin{align*}
\text{(25)} & \quad \text{if } a_{-a} d_A < ca_a p_A \\
& \quad \begin{cases} 
(A_0^0, A(1)) \text{ is of saddle type} \\
(A_0^0, A(2)) \text{ is a stable node}
\end{cases}
\end{align*}
\]

\[
\begin{align*}
\text{(26)} & \quad \text{if } a_{-a} d_A > ca_a p_A \\
& \quad \begin{cases} 
(A_0^0, A(1)) \text{ is a stable node} \\
(A_0^0, A(2)) \text{ is of saddle type}
\end{cases}
\end{align*}
\]

As, for \(c < a_{-a} d_A / a_a p_A\), \(A(2)\) is negative, the protein concentration at the steady state of the rate equations (24) is zero \((A(1) = 0)\). For \(c > a_{-a} d_A / a_a p_A\), the protein concentration at equilibrium is \(A(2) = \frac{cp_a a_a - d_A a_{-a}}{d_A a_a}\). Therefore, the conservation law (23) tune a bifurcation for \(c = a_{-a} d_A / a_a p_A\) (transcritical bifurcation), implying the existence of a threshold effect tuned by the conservation law parameter.

In Figure 5, we show the dependence of the protein steady state on the total concentration of the gene.

5.2. **Spatial distribution and steady states.** We have focused on genetic regulatory models without specifying a spatial localization. In a genetic network
Figure 5. Dependence of the protein steady state on the total concentration of the gene. Below the bifurcation or threshold $c = a_{-a}d_A/a_apa$, the equilibrium value of protein concentration $A_{eq}$ is zero, while above it takes the value $A_{eq} = (p_Aca_a - d_Aa_{-a})/(d_Aa_a)$. The parameters are: $a_a = 1.0$, $a_{-a} = 0.1$, $p_A = 0.2$ and $d_A = 0.1$.

Describing some biological process, the initial concentration of proteins can significantly vary in space, across tissues. This is specifically true in developmental processes, where often proteins show a spatial distribution with very sharp slopes that play a basic role in the establishment of body plans of organisms. A well known case is the Drosophila segmentation, where variations on protein concentrations across the embryo induces protein patterning, [Driever and Nüsslein-Volhard 1988, Nüsslein-Volhard 1992]. One of such genetic regulatory networks is the one represented in Figure 1 [Alves and Dilão 2006].

To show that patterning can be explained by the non homogeneity of initial conditions of regulators across tissues, we analyze a genetic regulatory network for the production of a protein $B$, regulated by one activator $A$ and two repressor proteins $R$ and $S$, Figure 6. To simplify our analysis, we take the competitive case, where the activator and the repressors bind to the same binding site of the operon of protein $B$.

To simplify further, we assume that the spatial distribution of the proteins $A$, $R$ and $S$ are constants. Under these conditions and with the regulation model developed in the package GeneticNetworks, we obtain the system of linear rate
Figure 6. Genetic regulatory network for the production of protein $B$ with one activator $A$ and two repressors $R$ and $S$.

\begin{align*}
B' &= p_B B_A - d_B B \\
(B_A)' &= b_a A B_0 - b_{-a} B_A \\
(B_R)' &= b_r R B_0 - b_{-r} B_R \\
(B_S)' &= b_s S B_0 - b_{-s} B_S
\end{align*}

(27)

where $B = B(x,t)$, $B_0 = B_0(x,t)$, $B_A = B_A(x,t)$, $B_R = B_R(x,t)$, $B_S = B_S(x,t)$, $A = A(x)$, $R = R(x)$ and $S = S(x)$. These concentration variables are considered to be defined in a spatial one-dimensional bounded region of the real line ($x \in I \subseteq \mathbb{R}$). The following conservation law holds,

$$B_0(x,t) + B_A(x,t) + B_R(x,t) + B_S(x,t) = c(x)$$

where $c(x)$ is a constant, depending eventually of the spatial independent coordinate $x$. The system of rate equations (27) has one steady state with coordinates,

$$\bar{B} = \frac{p_B \bar{B}_A}{d_B}, \quad \bar{B}_A = \frac{c A b_r b_{-s} b_{-a}}{D}$$

$$\bar{B}_R = \frac{c R b_{-a} b_{-r} b_{-s}}{D}, \quad \bar{B}_S = \frac{c S b_{-a} b_{-r} b_{-s}}{D}$$

where,

$$D = b_{-a} b_{-r} b_{-s} + Ab_r b_{-s} + R b_{-a} b_{-r} + S b_{-a} b_{-r} b_{-s}$$
Choosing $b_a = b_r = b_s$, and $b_{-a} = b_{-r} = b_{-s}$, the steady state concentration of the protein $B$, is,

\[ B(x) = \frac{c(x)A(x)p_B}{(1 + A(x) + R(x) + S(x))d_B} \quad (28) \]

In Figure 7, we show the steady state concentration (28) of protein $B$, as a function of a spatial coordinate, $x \in [0, 1]$. We have considered the initial distributions $A(x) = 0.8$, $R(x) = 83e^{-7x}$, $S(x) = 83e^{-7(1-x)}$, $c(x) = 100$, and the parameter value $p_B/d_B = 1$. For this case, the steady distribution of protein $B$ is spiky, due to the inhibitory regulation of the repressors. We have analyzed the same genetic network of Figure 6 with a model with different binding sites for each regulator. The final result is similar with the one shown in Figure 7. This shows that the two model approaches in GeneticNetworks, with one binding site and with several binding sites in the operon, give similar qualitative results. When, the calibration and validation of models is not a problem, we can use the simplest one binding site regulation model in order to describe a given genetic regulatory network. The one-regulator site framework leads to simpler mathematical models.

The solution (28) shows that steady states can depend on the initial conditions of the regulators ($A(x)$, $R(x)$ and $S(x)$) and on the initial conditions of the catalytic agents ($c(x)$), showing that spatial patterning can be a direct consequence of the non homogeneity of initial conditions.

In this model, we have considered that the concentrations of $A$, $R$ and $S$ are constants, implying that the model equations (27) describe a system where $A$, $R$ and $S$ have a fast recovery time. This situation only occurs in thermodynamically open systems, as is the case of biological systems.
5.3. Cross-regulation in *Drosophila*. In *Drosophila* early development, some proteins as Bicoid (BCD) and Hunchback (HB) are translated from mRNA of maternal origin. Early in the first developmental stages of *Drosophila*, at cleavage stage 13, BCD and HB proteins form a stable gradient along the antero-posterior axis of the *Drosophila* embryo, Figure 8. This stable gradient has its origin in a diffusion process occurring in the syncytial blastoderm of the embryo, (Dilão and Muraro, 2009). At a latter stage, in the 14th cleavage stage, other proteins as Knirps (KNI) show segments characterized by spiky concentration patterns along the antero-posterior axis of the embryo.

We show now that the patterning HB and KNI proteins as observed at late cleavage stage 14 of the embryo of *Drosophila* is due to the non-homogeneity of protein distribution at an early developmental stage. For that, we have taken the genetic regulatory network of Figure 1 describing the genetic regulation of HB and KNI, and we have used the package *GeneticNetworks* to describe the production of proteins Hunchback and Knirps during the cleavage cycles 14 of the developing embryo of *Drosophila*. 

**Figure 7.** Steady states of proteins $B$, $R$ and $S$ for the genetic regulatory network of Figure 6. The steady state of protein $B$ shows a spiky profile, resulting from the inhibitory action of proteins $R$ and $S$. In this model, we have considered that the concentrations of $R$ and $S$ are constant in time and non-homogeneous in space. The activator protein $A$ has been considered constant along the spatial region.
Figure 8. Concentration of protein Hunchback (HB) at the end of cleavage cycle 13, and of Bicoid (BCD) and Tailless (TLL) proteins at the cleavage cycle 14, along the antero-posterior axis of the *Drosophila* embryo. The embryo length has been scaled from 0 to 100, and the vertical axis units are proportional to protein concentration. The continuous curves represent the mean distribution of the concentration of proteins calculated for the data of 954 embryos. The data has been taken from the FlyEx database (Poustelnikova *et al.*, 2004; Pisarev *et al.*, 2009, http://flyex.ams.sunysb.edu/flyex/).

At the end of cleavage cycle 13 and beginning of the 14th, BCD, HB and TLL proteins have a non homogeneous distribution along the embryo, Figure 8.

Hunchback and Knirps proteins are both activated by the maternally produced protein Bicoid, Figure 1, and they mutually repress each other. The protein Knirps is also repressed by the protein Tailless of maternal origin. Therefore, the genetic network model obtained with the package *GeneticNetworks* should lead to the experimental profiles of HB and KNI, as observed at cleavage cycle 14, Figure 9.

In Figure 9, we show the experimental profiles of HB and KNI at cleavage cycle 14, as well as a fit of the experimental data obtained with a model built with the software package *GeneticNetworks*. The model equations have 22 free parameters. To fit the experimental data with the mathematical model, we have assumed that the initial protein concentrations of BCD and TLL are constant over time, and we have assumed the option of several binding sites per operon. To find the numerical
values of the model parameters in order to calibrate the model, we have used an optimization techniques based on a genetic algorithm, (Dilão et al., 2009; Dilão and Muraro, 2009b). We have considered that, in the ordinary differential equations of the model, the time is also a free parameter. The protein profiles shown in Figure 9 are out of steady state pattern, obtained for the integration time $t = 805$ s.

**Figure 9.** The dots with error bars are the mean experimental profiles of the proteins HB and KNI at cleavage cycle 14, for several Drosophila embryos along the antero-posterior axis. The embryo length has been scaled from 0 to 100, and the vertical axis units are proportional to protein concentration. The lines show the prediction of the model profile for proteins HB and KNI, obtained with the software package GeneticNetworks. The mean distribution of the concentration of proteins has been calculated for the data of 954 embryos, taken from the FlyEx database (Poustelnikova et al., 2004; Pisarev et al., 2009, http://flyex.ams.sunysb.edu/flyex/).

The fitted curves in Figure 9 shows a very good agreement with the experimental data. HB fits well in the embryo length range [5%, 80%], and KNI fits well in the embryo length range [20%, 100%]. The quality of the fits has been measured by the penalized chi square test. For the two fits in Figure 9, we have obtained the reduced chi square values $\chi^2_{HB} = 0.57$ and $\chi^2_{KNI} = 0.68$. This validates the genetic regulatory network of Figure 1. This suggest that the anterior and posterior regions of the embryo are under the control of additional regulators.
6. Conclusion

We have presented a software tool to build mathematical models of genetic regulatory systems. The input of the package is the graph containing the list of transcriptional activators and repressors of the network. The software implements an approach based on the mass action law, on the operon model for regulation, and we have assumed in general that genes are always catalytic substances presented in any genetically controlled biological process.

Within this approach, the usual threshold concept in biology emerges as a bifurcation phenomenon of the model equations. These bifurcations are in general tuned by the conservation law constants of the model equations. Therefore, within this framework, thresholds are emergent phenomena and they result from the catalytic role of genes, (Tyson, 2007).

Another consequence of the modeling approach presented here is that positional information in developmental processes can be described by non-homogeneity in the spatial distribution of regulators, and not necessarily associated with other physical processes of transport and diffusion.

Finally, we have validated a genetic regulatory network for the production of Hunchback and KNI during the 14th cleavage stage of the embryo of Drosophila, and we have presented evidence that gap gene protein segments are out of equilibrium patterns. The genetic regulatory network of Figure 1 describes well the gap gene protein concentration of HB in the embryo length region [5%, 80%], as well as the gap gene protein concentration of KNI in the embryo length region [20%, 100%].
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