

# Modeling Cell Populations in Development using Individual Stochastic Regulatory Networks

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**Abstract:** We present a new approach to high level stochastic simulations of cell populations. The proposed method employs the Stochastic Logical Network (SLN) method for simulating independent regulatory processes occurring in individual cells allowing for efficient simulations of systems consisting of thousands of cells. The stochastic logical network model is extended to account for not only regulatory control of gene expression but other related processes such as: inter-cellular signaling, cell division and programmed cell death. In the paper, we present the method and several case studies, where the proposed approach is used to provide models of biological phenomena. These examples include community effect in gene expression, the role of negative feedback in growing epithelial cell lineage and the role of asymmetric cell division in cell fate choices. We present also an efficient implementation of the method using GPU computing and show that its performance is significantly better than that using CPU.

## 1 INTRODUCTION

Modeling biological processes linked to gene regulation is a very broad and diverse field. The problem is fundamental and many different approaches with different advantages and disadvantages have been proposed to address its various aspects (see (De Jong, 2002) for a review). In particular, development of multi-cellular organisms presents many challenges as it requires consistent models taking into account large populations of cells able to perform their regulatory programs independently but at the same time able to communicate with each other via chemical signaling. In addition, the developmental systems need to be robust to external conditions and at the same time, they rarely operate under equilibrium conditions, requiring models capable of capturing temporal behavior as well as analysis of possible attractors and steady states of the system. In addition, frequently possible states of the system differ only by a small number of molecules leading to stochastic behavior on a single-cell level.

Boolean networks have been particularly useful in modeling developmental systems (Thomas and D'Ari, 1990), as they do not require exact rate constants and they provide the modelers with models which are relatively easy to study. However, most of the systems are focusing on cell autonomous be-

havior, with notable exceptions of several examples of networks related to *Drosophila* segmentation (Sánchez et al., 2008) or wing disc (Gonzalez et al., 2008), however even those pioneering works have only considered at most a handful of cells at once.

## 2 PROPOSED MODEL

We propose a new approach to modeling large cell populations with explicitly simulating regulatory networks of each of the cells. The method is based on stochastic logical networks (SLN), a generalization of the Boolean network model which builds up upon the non-deterministic state transitions in Boolean networks and defines a probabilistic model of state transitions. In the following sections we will briefly describe the SLN framework and how we use it for the purpose of simulating cell populations.

### 2.1 From Boolean Networks to Stochastic Logical Networks

Boolean networks have a long history of applications to gene regulatory systems. Early work by Kauffman (Kauffman, 1977) and Thomas (Thomas, 1978) led to the representation of the regulatory network as a sys-

tem of Boolean equations. For example a simple negative feedback loop can be represented by two equations:

$$\begin{aligned} X &\leftarrow Y \\ Y &\leftarrow -X \end{aligned}$$

In those first simple models of regulatory networks the dynamics was usually considered to be synchronous, i.e. all genes changed their value to their respective regulatory function (right-hand side of the regulatory equation). While the simplicity of such an approach might be appealing, it is now well known that due to very different transcription and decay rate for different genes, it is quite far from realistic.

Later, Thomas extended this formalism and proposed generalized logical networks (Thomas and D'Ari, 1990), which not-only introduced asynchronous state change for different genes, but also extended the state space to include more than two expression levels for some of the genes. The asynchronous state change had a major impact on the analysis of the system dynamics, as now instead of a single successor state for any state of the network we had to consider a set of possible successor states. While it is clear that the asynchronous state transitions are more realistic than synchronous, they also create challenges for the formal analysis of such systems as the size of the state space visited from any initial state can be much larger than in the synchronous case.

More recently, we proposed another extension of the Generalized logic formalism, by introducing stochastic models that could be built on top of the generalized logical networks. Stochastic logical networks (SLN) (Wilczynski and Tiurnyn, 2006) were originally defined as continuous dynamical systems with a canonical discretized form aimed at network reconstruction from quantitative data (Wilczynski and Tiurnyn, 2007), however for the purposes of this article, we will consider them to be discrete objects. Similarly to the Boolean models, any SLN with  $N$  genes consists of  $N$  variables  $g_1, \dots, g_N$  each of which takes values from a finite discrete set. For the purposes of this article, we will consider only variables with binary value sets, but all the reasoning can be extended to use variables with integer values and the proposed implementation is ready to use such extended variables. The regulatory function of each gene  $g_i$  is described by an equation of the following form:

$$F_i(g_1, \dots, g_N) = \sum_{j=1}^N w_{ji} g_j, \quad (1)$$

where  $w_{ji}$  is the regulatory influence of the gene  $g_j$  on  $g_i$ . For any state of the network, we can calculate the regulatory function for each of the genes

$\mathcal{F}_i = \{F_1, \dots, F_N\}$ . Given the matrix  $W = \{w_{ij}\}$ , SLN model defines a probability distribution

$$\mathcal{R}_i(\sigma) = \frac{|F_i|}{\sum |F_j|}$$

describing the probability of changing first the value of gene  $i$ , given that we start from the state  $\sigma$ . Given this probability distribution, we can employ an algorithm, conceptually similar to the Gillespie's stochastic simulation method (Gillespie, 1977) to simulate a cell behavior over time: in each step we select a gene to change based on its total influence  $F_i$  in comparison to the total influence in the system.

## 2.2 Modeling Populations of Cells with Stochastic Logical Networks

Starting with this slightly simplified definition of the SLN model, we can extend it to systems consisting of multiple cells with identical underlying genetic network. To achieve this, instead of a single state  $\sigma$ , we need to consider a population matrix  $P_{ij}$ , where the  $i$ -th row  $P[i]$  corresponds to a state of a single  $i$ -th cell.

With this representation, we can calculate the total influence of all genes by simply performing matrix multiplication of  $P$  by the regulatory matrix  $W$ . Once we have the influence, we just need to perform a single step of the simulation for each of the cells in the population to obtain the state of the population for the next step.

To account for typical processes involved in gene regulation in development, we have included additional actions performed at each step of population simulation:

- There are special variables, regulated in the same way as the gene variables, corresponding to the cell division, and death. After calculating the state of each cell, the algorithm checks whether the division or death variables are "active" and if it is the case, it either removes the cell from population or duplicates its row in case of symmetric division.
- For modeling signaling, a special vector is defined for mapping each of the genes to secreted molecules. Each secreted molecule is described by a positive integer. If such value is placed on the  $i$ -th position of the secretion vector and the  $i$ -th gene is active, a predefined number of molecules of the specified type is secreted to the environment. In case of non-secreted genes, the secretion vector is set to 0.
- After the secretion is performed, another vector mapping the signaling molecules back to genes is

used. If the number of molecules of a given signal is non-zero, for each cell capable of receiving such signal an integration event occurs randomly, with the probability proportional to the number of molecules divided by the number of cells in population.

This leads to the following algorithm:

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Algorithm 1: Algorithm for a single step of STOPS simulation.

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**Require:**  $W$  : regulatory matrix  
**Require:**  $P$  : population matrix  
 $F \leftarrow P \times W$   
**for**  $i=1$  **to**  $N$  **do**  
    choose a random  $j$  according to distribution  $F_i$   
    update  $P[i,j]$   
**end for**  
**for**  $i=1$  **to**  $N$  **do**  
    update Environment based on secretion from  $P[i]$   
**end for**  
**for**  $i=1$  **to**  $N$  **do**  
    update  $P[i]$  based on absorption from Environment  
**end for**  
**for**  $i=1$  **to**  $N$  **do**  
    multiply or delete row  $P[i]$  based on special variables  
**end for**

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Using this algorithm, we can simulate any SLN system provided that the matrix  $W$ , and the initial population are specified. We have written a prototype implementation of this method called STOPS (Stochastic Population Simulation) which is publicly available on-line and was used to obtain the results presented in the following section.

### 3 CASE STUDIES

In this section we describe three simple models of small biological systems representative to questions posed recently in the field of modeling regulatory networks in development. Each of those examples illustrates different capabilities of the STOPS modeling framework. All presented examples were implemented in STOPS and are publicly available for download (see Sec. 6).

#### 3.1 Community Effect in Gene Expression

Recent study by Saka *et al.* (Saka et al., 2011) proposed a simple model for the community effect in gene expression occurring during embryonic development of *Xenopus* frogs. During development, when it is necessary to create continuous tissues in the embryo it is usually achieved by formation of local aggregates of cells with correlated expression of a certain gene by means of signaling. The community effect definition is based on the observation that while such aggregates usually form around small foci of cells initially expressing the identity factor of the desired tissue, a certain number of cells capable of expressing this factor is required within the community to achieve stable activation of the whole colony. It was verified by dissection experiments, that if the number of cells is below a certain threshold, the activation of all cells is unlikely, while above this threshold the activation is prevalent among all cells.

This system can be represented by a simple gene network with signaling (see Fig. 1) as proposed by Saka *et al.* (Saka et al., 2011). It includes the identity gene  $g_1$ , which is responsible for production of the signaling molecule  $S$ , which is in turn secreted to the environment and from there can be sensed by all other cells via the receptor gene  $g_0$  capable in turn of activating the identity gene  $g_0$ . The whole network forms a simple positive feedback loop responsible for amplifying the initial signal, while the degradation rate of the signaling molecule is responsible for the threshold number of cells required for activation of majority of cells.

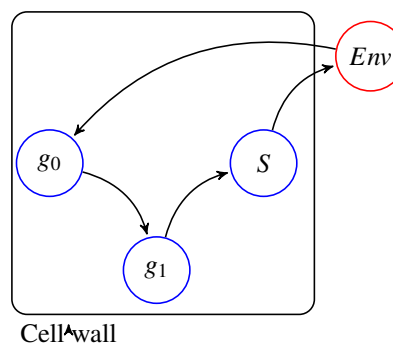


Figure 1: Simple gene regulatory network modeling community effect.

We have implemented this network in the STOPS framework and performed one hundred simulations, consisting of 30 steps each, for different colony sizes ranging from 50 to 300 cells. Then we measured for each of the trajectories, how many cells have the iden-

tivity gene ( $g_1$ ) active as a fraction of the whole population. For all cases, after 30 generations, we obtained cell colony either fully activated or fully silent, which is expected, given the presence of positive feedback in the system. It was also reassuring to see that the fraction of simulations leading to a fully activated system is exhibiting a stepwise dependence on the colony size: up to a 100 cells the silent case is clearly dominant, while starting from 150 cells the active state is dominating the results (see Fig. 2).

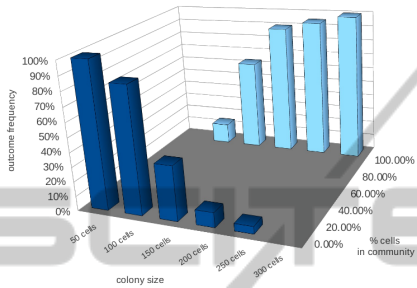


Figure 2: Outcomes of simulations of the community effect model.

### 3.2 Linear Cell Lineage with Proliferation

In a study by Lander *et al.* (Lander et al., 2009) the authors consider a simple scenario where there exists a pool of undifferentiated precursor cell population (expressing gene  $g_0$ ) which can then spontaneously switch to an intermediate differentiating state (expressing  $g_1$ ) which then leads to the fully differentiated terminal state (expressing  $g_2$ ). Importantly, only the undifferentiated states (expressing  $g_0$  or  $g_1$ ) can divide, giving rise to new undifferentiated cells, while the terminally differentiated cells (expressing  $g_2$ ) can enter apoptosis (programmed cell death). The system aims to model neural epithelia which consist of similar types of cells, which are important for their ability to quickly recover from removal of large numbers of differentiated and intermediate cells, provided that they are left with enough pluripotent cells. Using an ODE model for this system, Lander *et al.* observed that in order to achieve short recovery times and limited total number of cells it is beneficial to provide negative feedback from the differentiated cells instructing the pluripotent cells to limit their growth rate when the number of differentiated cells is sufficient.

Similar behavior can be observed in a system of cells modeled with Stochastic Logical Networks following the genetic regulatory interactions depicted in Figure 3. Each pair of genes representing consec-

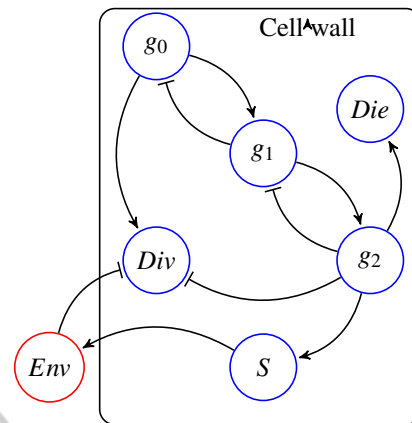


Figure 3: Gene network describing cell lineage with proliferation and signaling.

utive cell stages is linked by direct negative feedback loop providing basis for cell progression through the stages. There are two additional variables, corresponding to cell death and division and a signaling molecule released by the terminally differentiated cells and able to limit the proliferation of undifferentiated cells receiving the signal. Gene  $g_2$  corresponding to the terminal differentiation is also activating cell death and repressing proliferation.

To test whether the feedback cycle involving signaling is required for the functioning of the system, we can consider a system that is identical to the original one, but does not include the signaling component. Several exemplary trajectories of such a system are shown in Figure 4. While the general behavior of the system is consistent with earlier ODE simulations by Lander and colleagues, it should be noted, that the slope of recovery of the differentiated cells (blue lines) is not as steep as it should be and that the required number of intermediate cells (green line reaching 0.6 of the total population) are not supported by the experimental data. It is also disturbing to see that overall the population size is rapidly decreasing as a result of very quick removal of the primary cells from the population. It should be noted, that the problems of the sub-population of proliferating cells extinction is not visible in the original ODE model, as the pool of non-differentiated cells can get arbitrarily close to 0 and still be able to regenerate, while our stochastic model is capable of highlighting problems with models that lead to depletion of any sub-population as in our model it is possible to consider parameters leading to a complete extinction of the system.

If we compare these results to the full model including signaling to reduce proliferation when not needed, we can see clear improvement (see Fig. 4 B). The number of intermediate cells stays below 30



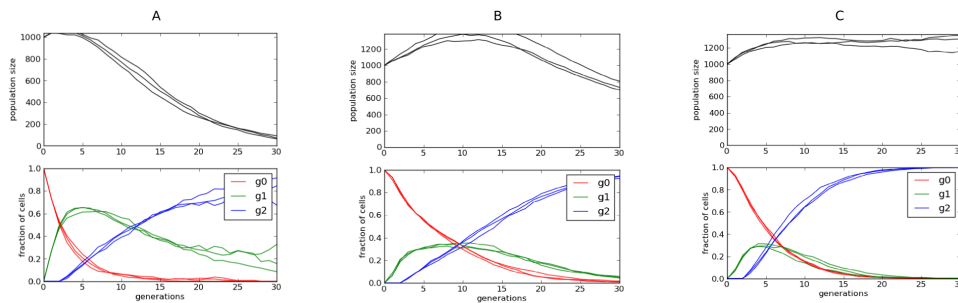


Figure 4: Results of simulation of the cell lineage without feedback (A), with simple feedback (B) and with cell-type specific feedback (C).

per cent all the time and the decrease in the number of primary cells is slower, however the regeneration rate of the differentiated cells remains unchanged and the population size problem is only slightly mitigated. So while it is clear that including feedback in the system improves the model, it is not sufficient to recapitulate the experimental results (faster regeneration and stable population size). It turns out, that the reason for these problems is the decreasing number of primary cells and the difficulty of the ligand to find those cells before being absorbed by the differentiated cells that cannot respond to the signal as they are already unable to proliferate. This problem cannot be seen in the ODE model, as there is dependence of the signal reception by the relative proportion of dividing cells in the population.

This problem can be eliminated by preventing the differentiated cells from being able to receive signals from the environment. Technically this can be done by including another gene representing the receptor, which is regulated by  $g_2$  and which is necessary for signal reception. Making this change to the system gives much better results (see Fig. 4 C). Both problems: slow regeneration rate and population size instability are solved making the model capable of reproducing experimentally observed behavior.

### 3.3 Asymmetric Cell Division and Cell-fate Choice

While signaling is a very powerful mechanism of controlling state changes in developing cells, it is frequently the case that the cell fate choice is determined by other mechanisms. Cohen *et al.* (Cohen et al., 2010) studied recently a system of neural cells differentiation, which includes two such mechanisms: asymmetric cell division and spontaneous stochastic cell-fate choice based on a bistable system of two genes.

The asymmetric division is a mechanism in which a cell expressing a pluripotency factor (in this case

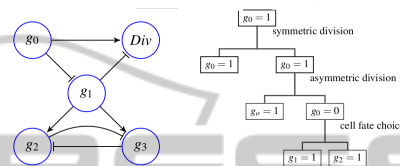


Figure 5: Network model of cell-fate choice with asymmetric cell division and the associated cell lineage tree.

a gene called “numb”) is undergoing an asymmetric division resulting with one of the daughter cells retaining majority of the factor protein, while the other daughter cell is born free of this protein. This leads these two cells to reach different regulatory states, since the pluripotency factor is lost in one of the resulting cells allowing it to start the differentiation process. On the regulatory level, the pluripotency factor is usually a repressor of the differentiation factors. In case of numb, it represses notch which is responsible for facilitating specification of cells into neurons (Shen et al., 2002).

The stochastic cell-fate choice facilitated by a bistable switch is a different regulatory mechanism which allows cells that already started differentiation to commit stably to one of the predefined cell-fates. It is nicely illustrated by the gene regulatory network for *Arabidopsis* flower development (Espinosa-Soto et al., 2004), where undifferentiated cells need to commit to one of the four cell-fates to make all necessary parts of the flower. On the regulatory level, such system, in its most basic form, consists of an initial signal: the differentiation factor which drives the expression of two, mutually exclusive, terminal cell-fate genes. Both these genes are repressing each other and repressing the upstream signal leading to the system with two attractors with either one (and only one) of the cell-fate genes on.

The system studied by Cohen and colleagues consisted of both those components: first it had undifferentiated cells capable of either self-reproduction by symmetric divisions or performing an asymmetric division leading to creation of a differentiating cell

which could then choose one of the two predefined cell fates. Interestingly, the authors provided experimental measurements of relative number of events leading to the symmetric vs. asymmetric divisions (53 vs. 19 respectively) and of the relative frequencies of choosing different cell-fates (10 vs 16). This allowed us to attempt to build a model capable of reproducing this behavior using the STOPS framework.

The regulatory network we designed is presented in Figure 5. It consists of the pluripotency factor  $g_0$  repressing the differentiation factor  $g_1$ , which has a basal expression level driving its expression immediately upon loss of  $g_0$ . Naturally, the proliferation factor activates the capability of cells to proliferate, while the differentiation gene is shutting this program down upon activation. The asymmetric division is acting on the expression level of  $g_0$ : assuming that the parent cell had the gene on an asymmetric division results with two cells which are identical to the parent cell regarding all genes but  $g_0$  which will be on in one of the daughters and off in the other.

Once the differentiation factor  $g_1$  is turned on, it starts driving the expression of both cell fate genes, however it is not necessarily acting on both of them with the same regulatory influence. In fact making those influences different is essential for reproducing the correct ratio between the numbers of different cell-fates in resulting population. At some point, one of the target gene gets activated subsequently leading to stable repression of the alternative cell fate. The overall cell-fate choice diagram is shown in Figure 5 (B).

The results of the simulation are capturing the general behavior of the system and by choosing the regulatory influences carefully, we were able to obtain simulation results matching the experimental results published by Cohen *et al.* (Cohen *et al.*, 2010).

## 4 IMPLEMENTATION AND PERFORMANCE BENCHMARKS

The first prototype of our stochastic population simulation (STOPS) method was implemented in pure python scripting language. While it was convenient for the prototype, the performance of such a solution was far from satisfactory. Since this tool is intended to be used for simulation of cellular populations with realistic sizes it needs to be able to tackle meaningful time-scales (thousands of simulation steps) for populations consisting of millions of cells. As we can see in Figure 6, the prototype implementation requires be-

tween 10 and 15 minutes for a single simulation step for a population of 10 million cells.

In order to provide a more efficient platform for realistic simulations, we have re-implemented the main algorithm in two different libraries dedicated to matrix operations: NumPy (Harrington and Goldsmith, 2009) and pyOpenCL (Klöckner *et al.*, 2012). The first one has the advantage of being available on all major platforms, making it possible for anyone to install STOPS on their machine and test it with reasonable efficiency. The second implementation uses the OpenCL bindings and allows users equipped with a machine with a support for an OpenCL implementation (currently there are OpenCL implementations released for AMD and Intel CPUs as well as Nvidia and ATI GPUs) to take advantage of the full potential of their hardware. We have tested the performance of all three implementations on a computer equipped with an Intel Xeon 3.2Ghz CPU and an Nvidia Tesla C1070 GPU. It is clear from Figure 6 that while all three implementations taking advantage of specialized matrix algebra operations outperform the initial version by an order of magnitude, it should be noted that the OpenCL version is still an order of magnitude faster than the NumPy version. Interestingly, the OpenCL performs similarly well both on CPU and GPU implementations of the OpenCL library. It is important, as the CPU has typically access to much larger memory, allowing for simulations of much larger systems. It should be noted that the memory consumption is the same (number of cells times the number of variables times the size of an integer).

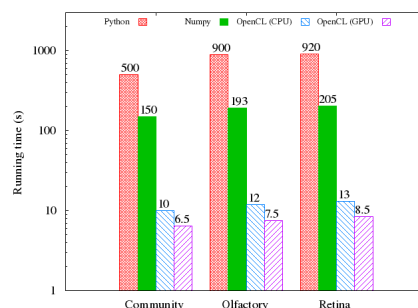


Figure 6: Comparison of running times for different STOPS implementations on three case-study datasets.

## 5 DISCUSSION AND FUTURE WORK

We presented here a new approach to modeling cell populations using stochastic logical networks. It is particularly well suited for developmental sys-

tems, where stochastic behavior of large populations of proliferating and signaling cells is driven by the same underlying regulatory machinery encoded in the genome. We also provide a prototype implementation capable of simulating cell populations with millions of cells on a standard personal computer. Even though the method is rather simple and requires only a handful of parameters to run a simulation, it is able to reproduce the results of many more established methods for wide variety of models relevant for problems currently under consideration by the modeling community. In some cases, like the linear cell lineage system, it can give us new insights missed by the ODE model due to its more accurate representation of small cell populations.

While the results shown are promising, the implementation is still in an early phase and could greatly benefit from multiple improvements. One key area that will work on in the future is extending the model to take into account spatial aspect of cell populations. Such functionality would greatly expand the range of possible applications of this model, however modeling spatially variable signalling without great decrease in the method performance poses a considerable challenge.

## 6 AVAILABILITY

The STOPS (STOchastic Population Simulation) software implementation is publicly available under the GNU GPL v.2 license. The implementation of all three case studies is included in current version available at <http://launchpad.net/stops>.

## ACKNOWLEDGEMENTS

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