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# Cell type Analysis from scRna-seq Data Achieved from a Mixture MOdel (CARDAMOM): vignette

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CARDAMOM (Cell type Analysis from scRna-seq Data achieved from a Mixture MOdel), is an algorithm for inferring a GRN from timestamped scRNA-seq data, which crucially exploits the notions of metastability and transcriptional bursting. It has been developed in Ventre. [1], and benchmarked and applied to real data in Ventre et al. [2]. In the latter, it has been shown that combined with the simulation algorithm HARISSA [3], the method allows to calibrate a mechanistic model in order to reproduce accurately experimental timestamped scRNA-seq data.

The aim of this vignette is to present the main functions of the method CARDAMOM, as well as to detail the small improvements that have been performed in Ventre, Herbach et al. [2] with respect to the first publication [1]. The actual package is available at the following address <https://github.com/eliasventre/cardamom>, and the previous version at <https://gitbio.ens-lyon.fr/eventr01/cardamom>.

## Brief description of the model

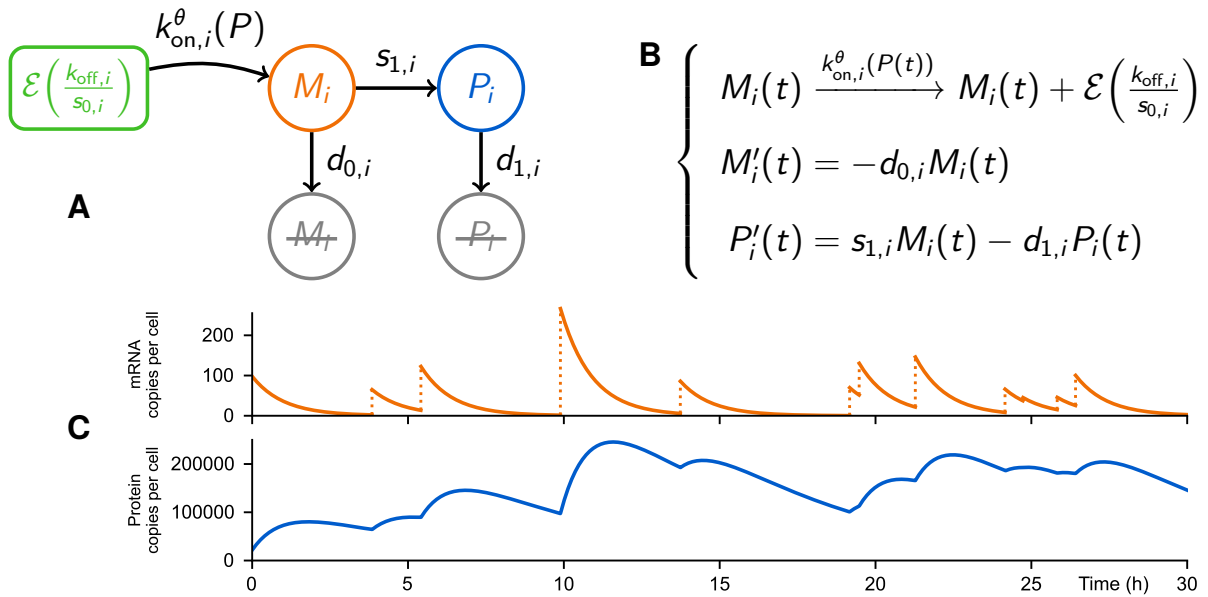
*This paragraph can be found on the Method section of [2].*

The model used throughout this article is based on a hybrid version of the well-established two-state model of gene expression [4], where a gene is described by the state of a promoter, which can be either *on* or *off*. If the promoter is *on*, mRNAs are being transcribed at a rate  $s_0$ , which are then translated into proteins at a rate  $s_1$ . Degradation of both mRNAs and proteins occurs at a rate  $d_0$  and  $d_1$ , respectively. The transitions between the on and off states occur at times of rates  $k_{on}$  and  $k_{off}$ . We consider the *bursty* regime of this model ( $k_{on} \ll k_{off}$ ), corresponding to short active periods with high transcription rates, as experimentally observed [5–8]. In this regime, mRNA is then transcribed by bursts of tens to hundreds of molecules. The random times at which these bursts occur are still described by an exponential distribution of parameter  $k_{on}$ , and their random size by an exponential distribution with mean  $s_0/k_{off}$ . This model is compatible with experimental single-cell data, as steady-state mRNA levels follow for each gene a Gamma distribution, in line with continuous single-cell data [9].

The key idea is to incorporate this model into a network: the burst rate for each gene  $i$  is given by a gene-specific function  $k_{\text{on},i}^\theta(P)$ , where  $P$  is the vector of protein quantities (Figure 1). This function depends on proteins through a GRN, represented by an  $n$ -by- $n$  matrix  $\theta = (\theta_{ij})$  where  $n$  is the number of genes in the network. The value of  $k_{\text{on},i}^\theta(P)$  then corresponds to the transcriptional burst frequency of gene  $i$  given protein levels  $P$ . Each parameter  $\theta_{ij}$  encodes the interaction  $j \rightarrow i$  with its direction, sign, and intensity. Recent work suggests that burst sizes are smaller and more uniform than previously anticipated [7] therefore leaving more room for burst frequency modulation [10] as a mechanism for gene expression regulation. We therefore consider that interactions come mainly from the modulation of burst frequencies  $k_{\text{on},i}^\theta$  and that for any gene  $i$ , the rates  $k_{\text{off},i}$  do not depend on  $P$ . The burst frequencies can be represented by sigmoid functions [3] as a simplification of the mechanistic form used in [4, 11]:

$$k_{\text{on},i}^\theta(P) = k_{0,i} + (k_{1,i} - k_{0,i}) \left( 1 + \exp \left( -\beta_i - \sum_{j=1}^n \theta_{ij} P_j \right) \right)^{-1} \quad (1)$$

where  $k_{0,i}$  (resp.  $k_{1,i}$ ) is the minimal (resp. maximal) burst frequency of gene  $i$  and  $\beta_i$  is the basal activity of gene  $i$ , which can be also considered as the constant activity of a set of genes that are not measured but act on the network.



**Figure 1:** Figure S1 from [2]. Graphical (A) and mathematical (B) descriptions of the mechanistic model for the dynamics of a gene  $i$ . (C) Bursts of mRNA occur at random times with rate  $k_{\text{on},i}$  and their size follows an exponential distribution  $\mathcal{E}(k_{\text{off},i}/s_{0,i})$ . The variables  $M_i$  and  $P_i$  describe respectively the mRNA and protein quantities associated to gene  $i$  in the cell. The vector of protein levels is denoted by  $P = (P_1, \dots, P_n)$  while  $\theta$  denotes the GRN which couples the genes together through functions  $k_{\text{on},i}$ .

## Description of the model parameters and their role in the dynamics

We provide here a list of the main variables of the mechanistic model. *It can be found on the Appendix A. of [1].*

- $\theta$  a matrix defining the interactions between genes, corresponding to a matrix with diagonal terms defining external stimuli,
- $k_{0,i}$  is the basal rate of expression of gene  $i$ ,
- $k_{1,i}$  is the maximal rate of expression of gene  $i$ ,
- $\beta_i$  is the basal activity of gene  $i$ , which can be also considered as the constant activity of set of genes which are not measured and act on the network,
- $d_{0,i}$  is the degradation rates for mRNAs of gene  $i$ ,
- $d_{1,i}$  is the degradation rate for proteins of gene  $i$ ,
- $s_{0,i}$  is the creation rate for mRNAs of gene  $i$ ,
- $s_{1,i}$  is the creation rate for proteins of gene  $i$ ,
- $k_{off,i}$  is the exponential rate of switching from state **on** to state **off** for the promoter of gene  $i$ ,
- $k_{on,i}$  is a sigmoidal function depending on the global protein field  $P \in \Omega$ , defined in (1), which characterizes the exponential rate of switching from state **off** to state **on** for the promoter of gene  $i$ ,
- $c_i = \frac{k_{off,i}}{s_{0,i}}$  is the exponential rate at which mRNAs of gene  $i$  are created at every burst.

The following table precise the way these parameters are encoded in an object of the class NetworkModel in the package CARDAMOM.

Parameters	Variable in the object NetworkModel
$k_0/d_0$	self.a[0]
$k_1/d_0$	self.a[1]
$c$	self.a[2]
$d_0$	self.d[0]
$d_1$	self.d[1]
$\theta$	self.inter
$\beta$	self.basal

Note that the creation rates  $s$  are not directly implemented because  $s_0$  is contained in  $c$  and  $s_1$  can always be fixed to  $S1 = s_1 := cd_0d_1/k_1$  which is the value for which the proteins scale to 1 (see Ventre. [1] Section 4.2). As the proteins are not observed in practice, this does not change the mRNAs counts obtained when simulating the model.

It is worth noticing to remark that as CARDAMOM only infers the parameters  $k_0/d_0$  and  $k_1/d_0$ , instead of  $k_0$  and  $k_1$ , the degradation rates completely determines the scale of the evolution speed. Indeed, the two first rows of the vector *self.a* inferred by CARDAMOM corresponds to the values of  $k_0/d_0$  and  $k_1/d_0$ . Thus, for a given gene  $i$ , a small  $d_{0,i}$  will lead to decrease the parameters  $k_{0,i}$  and  $k_{1,i}$  used for the simulations in HARISSA, making the dynamics slower in terms of both creation and degradation of mRNAs. In an equivalent way, a small  $d_{1,i}$  will also decrease the parameter  $s_{1,i}$  used for the simulations in HARISSA, making the dynamics slower in terms of both creation and degradation of proteins.

## Description of CARDAMOM

The principle of CARDAMOM is based on a two-step procedure:

- In a first step, we find the set of parameters defining the mixture of negative binomial distributions that best fits the data. The only parameter that can vary over time is the mixture proportion parameter  $\mu_t$  (allowing to estimate, for each gene  $i$ , the mean burst size  $s_{0,i}/k_{\text{off},i}$  and the values  $k_{0,i}$  and  $k_{1,i}$  of the typical modes associated to the function  $k_{\text{on},i}$ ). Note that all model parameters are estimated except the degradation rates, which are constant for each gene and scale the dynamics of gene expression.
- In a second step, we calibrate the basal activity and interaction parameters  $\beta_i$  and  $\theta_{ij}$  in order to approximate this mixture distribution. The interaction parameters  $\theta_{ij}$  are then updated at each timepoint sequentially to match the mixture parameters.

## Organisation of CARDAMOM

The functions of CARDAMOM are splitted into two directories:

- The first one is located at **cardamom/model** and contains the file **base.py**, that contains itself all the functions associated to an object of the class `NetworkModel` for calibrating the mechanistic model. The scRNA-seq data are given as an input to the function **fit**.
- The second one is located at **cardamom/inference** and contains the files **kinetics.py** and **network.py**. **kinetics.py** contains the functions allowing to perform the first step of CARDAMOM (finding the mixture parameters), and **network.py** contains the functions allowing to perform the second step of CARDAMOM (finding the associated network).

The function **fit** then uses the two main functions of the second directory, which are **infer\_kinetics** and **inference\_optim**.

- **infer\_kinetics**: This function infer the parameters of a mixture of Negative Binomial. In the first version of CARDAMOM [1], it was achieved using a MCMC algorithm on all the data observed at each timepoint grouped together. Following a simpler idea developed in [3], this function now simply fits the parameters of a Negative Binomial on the data observed at each timepoint and for each gene separately, using a variational method. For every gene, we consider that the rates  $k_{0,i}$  and  $k_{1,i}$  correspond respectively to the minimal and maximal values of these inferred parameters, following the assumption that every gene reaches its maximal frequency during one of the observed timepoints.
- **inference\_optim**: This function returns the inferred network (basal + inter), as well as the time at which each edge has been detected with the strongest intensity, as detailed in [1], Section 4. We have slightly modified the way each we penalized each coefficient, introducing weighting between the diagonal coefficients and the rest. This has been calibrated by cross-validation on a set of reference networks, and should not be changed for new networks.

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