Reactmine: a search algorithm for inferring chemical reaction networks from time series data

Julien Martinelli\textsuperscript{1,2,3} Jeremy Grignard\textsuperscript{1,4} Sylvain Soliman\textsuperscript{1} Annabelle Ballesta\textsuperscript{2} François Fages\textsuperscript{1} ⋆

\textsuperscript{1}Inria Saclay, Lifeware Group, Palaiseau, 91120, France, \textsuperscript{2}INSERM U900, Institut Curie, Saint Cloud, France, MINES ParisTech, CBIO - Centre for Computational Biology, PSL Research University, Paris, France, \textsuperscript{3}Department of Computer Science, Aalto University, Espoo, Finland, \textsuperscript{4}Institut de recherches Servier, Suresnes, France

⋆To whom correspondence should be addressed.

Abstract

Motivation: Inferring chemical reaction networks (CRN) from time series data is a challenge encouraged by the growing availability of quantitative temporal data at the cellular level. This motivates the design of algorithms to infer the preponderant reactions between the molecular species observed in a given biochemical process, and help to build CRN model structure and kinetics. Existing ODE-based inference methods such as SINDy resort to least square regression combined with sparsity-enforcing penalization, such as Lasso. However, when the input time series are only available in wild type conditions in which all reactions are present, we observe that current methods fail to learn sparse models.

Results: We present Reactmine, a CRN learning algorithm which enforces sparsity by inferring reactions in a sequential fashion within a search tree of bounded depth, ranking the inferred reaction candidates according to the variance of their kinetics, and re-optimizing the CRN kinetic parameters on the whole trace in a final pass to rank the inferred CRN candidates. We first evaluate its performance on simulation data from a benchmark of hidden CRNs, together with algorithmic hyperparameter sensitivity analyses, and then on two sets of real experimental data: one from protein fluorescence videomicroscopy of cell cycle and circadian clock markers, and one from biomedical measurements of systemic circadian biomarkers possibly acting on clock gene expression in peripheral organs. We show that Reactmine succeeds both on simulation data by retrieving hidden CRNs where SINDy fails, and on the two real datasets by inferring reactions in agreement with previous studies.

Availability: https://gitlab.inria.fr/julmarti/crninf/
Contact: francois.fages@inria.fr
Supplementary information: S1 detailed results of Reactmine and SINDy

1 Introduction

With the automation of biological experiments and the increase of quality of cell measurements, automating the building of mechanistic models from data becomes conceivable and a necessity for many new applications. The structure of such models, e.g. gene regulatory networks (GRN) or chemical reaction networks (CRN), is classically built from an extensive review and compilation of the literature by the modeler. More recently, efforts have been made to develop model learning algorithms to assist modelers in order to partly automate the model building process, in particular when time series measurements are available.
Extensive literature is available in the context of GRN inference or unsupervised learning, partly motivated by knowledge discovery problems, such as presented in the DREAM series of challenges (Stolovitzky et al. 2007), or experiment design (King et al. 2004). A GRN consists in a directed graph $G = (W, E)$ of genes and edges $E_{ij}$ between genes, whenever a gene transcription factor $W_i$ binds to the promoter region of target gene $W_j$. GRN inference algorithms feature a wide range of machine learning methods, e.g. Decision Trees (Huynh-Thu and Geurts 2018), Information Theory (Zoppoli et al. 2010) or Gaussian Processes (Aalto et al. 2020).

Less work concerns CRN inference, i.e. the problem of inferring both the structure and kinetics of chemical reactions between some molecular species observed with time series data about their concentrations. The structure of a CRN can be represented by a bipartite directed graph with edges from molecular species vertices to reaction vertices, representing the reactants of a reaction, and edges from reaction vertices to species representing their product. Of note, the indegree and outdegree of a reaction node can be above one, which allows for bimolecular reactions like complexations, e.g. $A + B \rightarrow C$, or catalyzed transformations, e.g. $A + B \rightarrow A + C$. Each reaction of a CRN is given with its kinetics, using reaction rate functions such as mass action law, Michaelis-Menten or Hill kinetics. The rate function of each reaction appears as a term in the ordinary differential equations (ODE) that govern the time evolution of the products and reactants of the reaction. Overall, both the difference of structure and the importance of the kinetics make the above GRN inference methods hardly applicable to CRN inference problems.

CRN inference may thus rely on the ODE semantics of a CRN to apply ODE inference methods from time series data, such as the state-of-the-art tool SINDy (Sparse Identification of Nonlinear Dynamics, Brunton et al. 2016) with an appropriate library of kinetic functions. The main assumption is that the dynamics of each variable can be expressed using only a few functions of the observed variables, without introducing hidden variables, so that techniques like sparse regression can be used to determine the optimal members of the library for a given problem. Selecting the ground truth sparse set of predictors is however a task best achieved provided two hypotheses are satisfied: low correlations between the true predictors and the spurious ones, and low partial correlations among the set of true predictors (Zhao and Yu 2006). These conditions can reasonably be met in datasets composed of multiple initial states with various combinations of absent and present species, possibly obtained by silencing genes of interest (i.e. knockout experiments) or exposure to targeted inhibitors. Such datasets containing time series in multiple conditions indeed allow the different reactions to be witnessed in an independent manner (Carcano et al. 2017). However, this is not always possible, and in many situations, like in the context of experimental time series data obtained from protein fluorescence microscopy (Feillet et al. 2014), one has to work only with traces obtained in a wild type setting in which those hypotheses are not satisfied.

In this paper which extends (Martinelli et al. 2019), we present Reactmine, a bounded-depth tree search algorithm to infer CRNs from time series data, without any low correlation assumption. Sparsity is enforced by inferring reactions with their kinetics in a sequential fashion, with the depth of the search tree bounding the number of inferred reactions. At each node, the reaction candidates are ranked according to the variance of the kinetics inferred on their transition support, and the best candidates are used as choice points at that node. At each successor node, one selected reaction is added and its effect is subtracted from the trace. Each leaf in the search tree represents a CRN candidate. In a final pass, the kinetic parameters of the leaf CRNs are globally re-optimized on the whole trace transitions, and the CRN candidates are ranked according to the quadratic loss between the predicted and experimentally-observed temporal variations. As an example, Figure 4 shows the recovery by Reactmine of the chain CRN $A \rightarrow B \rightarrow C \rightarrow D \rightarrow E$ with mass action law kinetics and rate constants equal to one, from a single simulation trace of $A, B, C, D$ and $E$ with $A$ initially present.

On a benchmark of synthetic data obtained by simulation from a hidden CRN with standard initial conditions, we show the capability of Reactmine to recover either the hidden CRN, or a variant CRN capable of reproducing the simulation data in the same range of initial conditions, whereas SINDy fails to infer sparse ODE systems and even to reproduce the time series data on different initial conditions. In these examples, we analyze the sensitivity of the results to
the number of time points and to the four hyperparameters of our algorithm: $\gamma$, the maximum number of reactions inferred (i.e. maximum depth of the search tree), $\beta$, the maximum number of reaction candidates (i.e. maximum branching factor of the search tree), $\delta_{\text{max}}$, the velocity similarity threshold between species taking part in a given reaction and $\alpha$, the variation coefficient acceptance threshold about the inferred kinetics on the supporting transitions.

Then, we apply Reactmine on two sets of real experimental data: one from protein fluorescence videomicroscopy of cell cycle and circadian clock markers in mammalian fibroblasts, and one from biomedical measurements of systemic circadian biomarkers possibly acting on clock gene expression in peripheral organs. We show that Reactmine succeeds in inferring meaningful interactions, interestingly in accordance with the main conclusions drawn from previous analyses of these datasets though ODE models built, respectively, using a temporal logic approach in [Traynard et al. (2016)], and a different model learning approach in [Martinelli et al. (2021)].

The rest of the article is organized as follows. In the Methods section, we present our CRN inference algorithm, its theoretical complexity and comparison to related work. In the Results section, we first evaluate its performance on synthetic data obtained by simulation of some hidden CRNs, and perform the above-mentioned sensitivity analyses. Then we show the results obtained with simulation data from the MAPK signaling CRN studied in [Qiao et al. (2007)]. In all those instances, our results are shown to compare favorably to SINDy. Then, we present our results on the two real-world biological datasets of this study, and compare them to the previous models developed from those data. Finally, we conclude on the merits of this approach and its current limitations.
2 System and methods

2.1 Notations

Bold lower (resp. upper) case letters denote vectors (resp. matrices). Unless stated otherwise, sets are represented with capital letters. For a matrix $\mathbf{M}$, $\mathbf{M}_{l,\bullet}$ (resp. $\mathbf{M}_{\bullet,i}$) stands for its $l^{th}$ row (resp. $i^{th}$ column).

We observe a system $\mathbf{y}$ describing the evolution of $m$ biological species at $n$ discrete time points $\{t_l\}_{1 \leq l \leq n}$. We focus on the case of a single trace, containing observations for all species at a finite set of time points represented by a data matrix of the form:

$$
\mathbf{Y} =
\begin{bmatrix}
  y_{1,1} & y_{1,2} & \cdots & y_{1,m} \\
  y_{2,1} & y_{2,2} & \cdots & y_{2,m} \\
  \vdots & \vdots & \ddots & \vdots \\
  y_{n,1} & y_{n,2} & \cdots & y_{n,m}
\end{bmatrix}
$$

The matrix of observed velocities, $\mathbf{V} = (v_{l,i})_{1 \leq l \leq n, 1 \leq i \leq m} \in \mathbb{R}^{n \times m}$ can be estimated from original data and has the same number of rows as $\mathbf{Y}$. The extension to multiple traces is straightforward by concatenation of the data and velocity matrices.

2.2 Chemical Reaction Networks

A chemical reaction is formally defined as a triple $(R, P, f)$, where $R$ (resp. $P$) is a multiset of reactant (resp. product) species and $f : \mathbb{R}_{+}^{n} \rightarrow \mathbb{R}_{+}$ is a rate function over molecular concentrations specifying the reaction kinetics. A reaction catalyst is a species in $R \cap P$. A chemical reaction network (CRN) is a finite set of reactions.

For the sake of simplicity in this article, we shall restrict ourselves to reactions with 0/1 stoichiometry only, and shall consider the stoichiometry vector of a reaction, $s \in \{-1, 0, 1\}^m$ where $\forall i \in [1, m]$, $s_i = 1$ if $i \in P \setminus R$, $-1$ if $i \in R \setminus P$, 0 otherwise. This excludes autocatalysis reactions for instance, although such reaction schemas could be accommodated as well.

We consider three types of kinetics for the reactions: mass action law, Michaelis-Menten and Hill kinetics. For a reaction with mass action law kinetics, the reaction rate is the product of the reactant species concentrations multiplied by some rate constant $k$. Such a reaction will be written $R \rightarrow P$.

2.3 Reactmine algorithm

Reactmine is a bounded-depth tree search algorithm which, at each node of the search tree,  

1. infers new reaction candidates composed of reactants and products with highest and similar changes on some observed transitions called their support, 
2. ranks them according to the variance of the ratio between observed and inferred velocities on their supports, 
3. and selects the $\beta$ best reaction candidates to add as successors in the search tree, with appropriate updates of the velocity matrix.

At the end, a global re-optimization of the kinetic parameters of the inferred CRNs at the leaves of the search tree is performed on the whole trace, in order to select the inferred CRN that minimizes the quadratic loss between the inferred and experimentally observed species velocities on the data. Reactmine thus uses four hyperparameters:

- $\gamma$, the depth bound on the search tree, i.e. the maximum number of inferred reactions along a branch, 
- $\delta_{\max}$, the velocity similarity threshold between the reactants and products of one reaction candidate in one observed transition, 
- $\alpha$, the kinetics variance threshold of one reaction candidate on its supporting transitions,
• $\beta$, the maximal number of reaction candidates selected at a node.

The different phases of the algorithm are detailed below.

2.3.1 Generation of reaction skeletons

Let us denote by $r = (R, P)$ a reaction skeleton, i.e. a reaction without rate function. Let us consider $\hat{V}_{l, \star}$, the velocity of the system for an arbitrary time point $t_l$. For any $1 \leq l \leq n, 1 \leq i \leq m$, let

$$\hat{v}_{l,i} = \max_{1 \leq j \leq n} |\hat{v}_{l,j}|$$

be the velocity of species $i$ at time $l$ normalized to its maximum velocity measured on the whole trace. Let

$$i_{\max} = \arg \max_i |\hat{v}_{l,i}|$$

Species $i_{\max}$ has the highest normalized velocity at time $t_l$. For that reason, that species is assumed to undergo the primary change that should be explained by one preponderant reaction at that time point. To determine the other components of the reaction, the following sets of reactants and products are computed:

$$R_{\delta}(t_l) = \left\{ i \in \{1, \ldots, m\}, \hat{v}_{l,i} < 0, \left|\frac{\hat{v}_{l,i_{\max}}}{\hat{v}_{l,i}}\right| \leq \delta \right\}$$

$$P_{\delta}(t_l) = \left\{ i \in \{1, \ldots, m\}, \hat{v}_{l,i} > 0, \left|\frac{\hat{v}_{l,i_{\max}}}{\hat{v}_{l,i}}\right| \leq \delta \right\}$$

Let $r_{\delta}(t_l) = (R_{\delta}(t_l), P_{\delta}(t_l))$ be the corresponding candidate reaction skeleton. Elements belonging to $R_{\delta}(t_l)$ or $P_{\delta}(t_l)$ have similar absolute variations compared to species $i_{\max}$. The upper bound of this sequence of $\delta$ values is the hyperparameter $\delta_{\max}$ of the algorithm. Equation (3) shows that it stands for the maximum absolute fold change allowed between the variations of species involved in a reaction. A value significantly above 1 accounts for the fact that $\hat{v}_{l,i_{\max}}$ might not be completely explained by only one reaction. The computation of $r_{\delta}(t_l)$ is therefore performed for various $\delta$ values as well as for all time points $\{t_l\}_{1 \leq l \leq n}$.

Now, let us define the support $T(r)$ of a reaction skeleton $r = (R, P)$, as the set of time points indices where it has been witnessed, e.g. in Figure 1:

$$T(r) = \{ l \in \{1, \ldots, n\}, \exists \delta \in [1, \delta_{\max}], r_{\delta}(t_l) = (R, P) \}$$

2.3.2 Inference of reaction kinetics

The next step consists in assigning a rate function to the reaction skeleton, to completely define one reaction. $(R, P, f)$ follows the law of mass action with parameter $k$ if $\forall j \in R \cup P, \forall l \in \{1, \ldots, n\}$

$$v_{l,j} = s_j f(Y_{l, \star}) = s_j k \prod_{u \in R} y_{l,u}$$

where we recall that $s_j$ is the stoichiometry of species $j$ in the reaction. Using the finite differences estimate $\hat{V}$ as well as the support set $T(r)$ for the current reaction candidate $r = (R, P)$, one can provide an estimator of $k \forall j \in R \cup P$:
\[ \hat{k}_j = \frac{s_j}{\# \mathcal{T}(r)} \sum_{l \in \mathcal{T}(r)} \frac{\hat{v}_{l,j}}{\prod_{u \in R} y_{l,u}} \]  

This estimator is designed to realize an equality in mean across the support between observed kinetics and inferred kinetics. The former is represented by the numerator, the latter by the denominator times \( \hat{k}_j \).

Reactmine is also compatible with other forms of kinetics, provided that a measure of quality of reaction parameters can be computed, such as the coefficient of variation. A single-reactant reaction \((R, P, f)\) follows Michaelis Menten kinetics if \( \forall j \in R \cup P, \forall l \in \{1, \ldots, n\} \)

\[ v_{l,j} = s_j f(Y_{l,*}) = s_j \nu_{\text{max}} \frac{y_{l,u}}{y_{l,u} + K_m} \]  

where \( \nu_{\text{max}} \) and \( K_m \) are parameters, and \( R = \{u\} \).

As \( y_u \to +\infty \), \( |v_j| \to \nu_{\text{max}} \). Besides, \( |v_j| \) being an increasing function of \( y_u \), an estimator of \( \nu_{\text{max}} \) can be obtained as the highest observed velocity \( v_{l,j} \) on the whole transitions: for all \( j \in R \cup P \)

\[ \tilde{\nu}_{\text{max},j} = \max_{l \in \{1, \ldots, n\}} |\hat{v}_{l,j}| \]  

Then, \( K_m \) is defined as the value of reactant concentration for which the associated velocity is equal to \( \frac{\nu_{\text{max}}}{2} \). Since measurements are only available at discrete time points, one has

\[ \hat{K}_{m,j} = y_{l^*,u} \quad \text{with} \quad l^* = \arg\min_{l \in \{1, \ldots, n\}} \left| \hat{v}_{l,j} - \frac{\tilde{\nu}_{\text{max},j}}{2} \right| \]  

Once an estimator for \( K_m \) has been provided, we apply the same principle as in Equation 7 to obtain a new estimator of \( \nu_{\text{max}} \) \( \forall j \in R \cup P \)

\[ \hat{\nu}_{\text{max},j} = \frac{s_j}{\# \mathcal{T}(r)} \sum_{l \in \mathcal{T}(r)} \hat{v}_{l,j} \hat{K}_{m,j} + y_{l,u} \]  

The computation described above also applies to Hill Kinetics of order \( \eta \), \( \forall j \in R \cup P, \forall l \in \{1, \ldots, n\} \)

\[ v_{l,j} = s_j f(Y_{l,*}) = s_j \nu_{\text{max}} \frac{y_{l,u}^\eta}{y_{l,u}^\eta + K_m} \]  

### 2.3.3 Ranking of the best reaction candidates

In order to compare reaction candidates between them, an interesting criterion to look at is certainly the statistical quality of the inferred kinetics on the support of the inferred reaction skeleton. The variance of the mass action law coefficient estimate over the support of the reaction can be estimated itself for each species involved in the reaction:

\[ \sigma_j = \frac{1}{\# \mathcal{T}(r)} \sum_{l \in \mathcal{T}(r)} \left( \frac{\hat{v}_{l,j}}{\prod_{u \in R} y_{l,u}} - \hat{k}_j \right)^2 \]  

It is worth noticing however that there is a relationship between mean and variance when estimating the kinetics of different reactions: a slow reaction will tend to produce a low variance, compared to a faster reaction. We thus consider the coefficient of variation (CV),

\[ \rho_j = \frac{\sigma_j}{|\hat{k}_j|} \]  

measured for each reactant or product of the reaction, and introduce more precisely the species index that minimizes it:

\[ j^* = \arg\min_{j \in R \cup P} \rho_j \]
on which we rely to estimate $k$. The complete reaction is therefore $r = (R, P, \hat{f})$ with $\hat{f} : y \mapsto \hat{k} \prod_{u \in R} y_u$
and $\hat{k} = k_{j^*}$. This process is performed for all reaction skeleton candidates.

A reaction candidate $r$ is accepted if it satisfies $\rho(r) < \alpha$. A typically acceptable value for $\alpha$ is below 1, indicating that the variance of the estimator does not overcome the mean. In the event where the best reaction $r^*$ fails to satisfy that condition, the addition of a catalyst to the reaction candidate is tried. To that end, Equation (6) is modified:

$$v_{l,j} = s_j \hat{k} \prod_{u \in R \cup \{c\}} y_{l,u}$$

(16)

$\forall j \in R \cup P$, and $c$ can be any species. The optimal catalyst $c^*$ for a particular reaction is the species providing the lowest CV, in which case $R \leftarrow c^*$ and $P \leftarrow c^*$. A catalyzed reaction is accepted if its associated loss value is below $\alpha$.

The reaction candidates are thus ranked according to their CV, the lowest CV corresponding to the best reaction. The $\beta$ best accepted reactions are returned, representing the maximum number of inferred candidates.

### 2.3.4 Velocity matrix update

Once a candidate reaction $(R, P, f)$ is accepted and selected to develop the search tree, its effect on the velocity data is removed at that node of the search tree as follows:

$$\hat{V} \leftarrow \hat{V} - \begin{pmatrix} f(Y_{1,*}) \\ \vdots \\ f(Y_{n,*}) \end{pmatrix} s^T$$

(17)

This mechanism is illustrated in Figure S1 with the example of the chain CRN.

It is worth remarking that this velocity update mechanism can also be used in our approach to take into account prior knowledge consisting of already known reactions between the observed species, simply by updating the initial velocity matrix according to those reactions.

### 2.3.5 Theoretical complexity

**Proposition.** The computational time complexity to infer one reaction $(R, P, f)$ is $O(nmI)$ where $n$ is the number of time points, $m$ the number of species, and $I = |R \cup P|$.

**Proof.** Inferring the reaction kinetics constant involves the computation of a mean for each species present in the reaction (Equation [7]), which is $O(nI)$. In the worst-case, a lookup for a catalyst species is necessary, at a cost of $O(nIm)$. The velocities update step performed in Equation [17] is $O(nI)$. Generating reaction skeletons requires the computation of the species displaying highest velocities for each time point, which is $O(nnm)$ (Equation [3]). After that, the sets $R_q(t_l)$ and $P_q(t_l)$ are obtained with a bounded number of $\delta$ values. The time complexity for the inference of one reaction is therefore $O(nnmI)$.

Since the depth of the search tree is bounded by $\gamma$ and each node has at most $\beta$ children, the time complexity of Reactmine is thus $O(\beta^\gamma nmI)$.

### 2.3.6 Final global re-optimization of kinetic parameters

The sequential inference of reactions together with their kinetics may introduce errors in the rate constants which can be corrected by a final global re-optimization step of the rate constants of all reactions on the whole trace. Let $V$ be the initial matrix of the estimated velocities, i.e. before the removal of the effect of the inferred reactions. Once a CRN has been built from the iterative inference of $p$ reactions, an additional optimization step can be applied. From a CRN $R = \{(R_q, P_q, f_q)\}_{1 \leq q \leq p}$ and data matrix $Y$, one can construct a matrix $F(Y, k) \in \mathbb{R}^{n \times p}$:

$$F(Y, k) := \begin{bmatrix} f_1(Y_{1,*}, k) & \ldots & f_q(Y_{1,*}, k) & \ldots & f_p(Y_{1,*}, k) \end{bmatrix}$$

(18)
with \(n\) the number of time points. The \(q\)th column of \(\mathbf{F}(\mathbf{Y}, \mathbf{k})\) is a vector describing the rate of the reaction \((R_q, P_q, f_q)\) at each time point, with \(\mathbf{k}\) being the vector of reaction kinetic parameters. Combined with the stoichiometry matrix of the CRN \(\mathbf{S} \in \mathbb{R}^{p \times m}\), we can formulate an optimization problem:

\[
\mathbf{k} = \arg\min_{\mathbf{k} \in \mathbb{R}_+^p} \| \hat{\mathbf{V}} - \mathbf{F}(\mathbf{Y}, \mathbf{k})\mathbf{S} \|^2_F
\]  

(19)

Where \(\| \cdot \|_F\) is the Frobenius norm. In particular, for a mass action law CRN,

\[
\mathbf{F}(\mathbf{Y}, \mathbf{k}) = \left[ \prod_{i \in R_1} y_{i,1} \cdots \prod_{i \in R_q} y_{i,i} \cdots \prod_{i \in R_p} y_{i,i} \right] \text{diag}(\mathbf{k})
\]  

(20)

The optimization starts with an initial guess set as \(\mathbf{k} = (k_1, \ldots, k_p)^T\). It is worth noticing that the least squares term compares the inferred and observed velocities, rather than the data measurements \(\mathbf{Y}\) and a numerical integration of the inferred CRN, which allows avoiding the resolution of a non-convex optimization problem in the case of mass action law kinetics. Indeed, Equation (20) shows that the inferred velocities are written as a weighted linear combination of reaction effects, which makes the minimization problem convex. However, for Michaelis-Menten and Hill kinetics, the optimization of \(\hat{\mathbf{V}}\) might span a wide range from one species to another, we normalize the convergence guarantees. Furthermore, in order to take into account the fact that concentrations and Hill kinetics, the optimization of \(K_m\) constants leads to a non-convex problem, with no convergence guarantees. Moreover, in order to take into account the fact that concentrations might span a wide range from one species to another, we normalize the \(q\)th column of the matrix \((\mathbf{V} - \mathbf{F}(\mathbf{Y}, \mathbf{k})\mathbf{S})\) by \(\max_{1 \leq l \leq n} \mathbf{V}_{l,j}\), for all \(j\), inducing equal importance for each species in the cost function. This final step of global optimization is performed for all CRN candidates at the leaves of the search tree. Reactmine returns the CRN which minimizes the loss function defined in Equation (19).

Finally, it is worth noting that the same loss function can be used more globally to parametrize Reactmine by choosing the hyperparameter values which minimize the loss. This is done by grid search as described in the next section on evaluation results.

### 2.4 Related work

Some methods originally designed to discover the dynamics of physical systems can be applied to our problem of inferring a CRN from time series data. Most notably, the SINDy system [Brunton et al. 2016], starting from temporal measurements, aims at providing a reconstruction of the velocities in the following way:

\[
\hat{\mathbf{V}} = \Theta(\mathbf{Y})\mathbf{Ξ}
\]  

(21)

\(\Theta(\mathbf{Y}) \in \mathbb{R}^{n \times p}\) is a library of \(p\) functions constructed from the input variables \(\mathbf{Y}\) including, for instance, first to \(m\)-order polynomial interactions, e.g. \(\mathbf{Y}_{\ast,1} \odot \mathbf{Y}_{\ast,j}\), the sin and cos functions, e.g. \(\sin(\mathbf{Y}_{\ast,j})\), or even more sophisticated user-defined functions. The dynamics of each variable is then captured by a weighted combination of library members, the weights being encompassed in \(\mathbf{Ξ}\). Because it is thought that the expression of the dynamics should be sparse within the library \(\Theta(\mathbf{Y})\), SINDy proposes to obtain \(\mathbf{Ξ}\) using sparse regression.

\[
\mathbf{Ξ} = \arg\min_{\mathbf{Ξ} \in \mathbb{R}^{p \times m}} \| \hat{\mathbf{V}} - \Theta(\mathbf{Y})\mathbf{Ξ} \|^2_F + \lambda \| \mathbf{Ξ} \|_1
\]  

(22)

For fair comparison within our CRN setup with mass action law and stoichiometry at most 1, \(\Theta(\mathbf{Y})\) is here restricted to polynomials up to the second order, and a bias term. Hence \(p = \left(1 + \frac{m(m+1)}{2}\right)\).

\[
\Theta(\mathbf{Y}) := \begin{bmatrix} 1 & \mathbf{Y}_{\ast,1} & \cdots & \mathbf{Y}_{\ast,m} & \mathbf{Y}_{\ast,1}\mathbf{Y}_{\ast,2} & \cdots & \mathbf{Y}_{\ast,m-1}\mathbf{Y}_{\ast,m} \end{bmatrix}
\]  

(23)

Associated to a positive weight, the bias term corresponds to a synthesis reaction. First order interactions translate to reactions such as \(A \xrightarrow{\Delta} B + C\) for \(A \neq \emptyset\), with the special case \(A \in \{B, C\}\).
corresponding to a catalyzed synthesis. Second order interactions encompass reactions of the form \( A + B \xrightarrow{k} C + D \) for \( \{A, B\} \neq \emptyset \). Again, the case \( A \neq B \in \{C, D\} \) corresponds to a catalyzed reaction, with \( Y \) referring to the exclusive OR.

SOMETHING RELATED TO SINDy is the algorithm GRISLI originally designed for inferring Gene Regulatory Networks from single-cell data (Aubin-Frankowski and Vert, 2020). GRISLI solves a similar problem as SINDy, but estimates velocities \( V \) thanks to a weighted average of finite differences, with weights defined by a spatio-temporal kernel \( K(Y_{t,i}, Y_{t,i'}) \), which quantifies how \( Y_{t,i} \) is believed to be useful in estimating the velocity at \( Y_{t,i'} \). Subsequently, multiple instances of Equation (22) are solved, each time with a bootstrapped sample of \( Y \) and \( V \). This leads to frequencies of apparition of each term. GRISLI considers a library \( \Theta(Y) \) made with first-order interactions only, which makes it a particular case of SINDy for the optimization part. This last point means that GRISLI cannot infer complexation reactions. Furthermore, GRISLI outputs frequencies that are difficult to compare with the output of Reactmine and SINDy. For these reasons, we decided to compare with SINDy only. In our experiments, we use the \texttt{pySINDy} package with STLSQ optimizer, as the latter yielded the best results.

\section{Results}

\subsection{Evaluation on simulation data from hidden CRNs}

We first evaluate our algorithm on simulation data obtained from hidden CRNs, with which the inferred CRNs can be compared. For Reactmine, we report the reactions in the order of their inference along the search tree branch that gives the best CRN. For SINDy, we report the inferred ODE systems. In these experiments, numerical integration is performed using the Python package \texttt{scipy.integrate.odeint} with the default integrator \texttt{lsoda}. Simulations run for a time horizon \( T = 10 \) and a time step \( \Delta t = 0.1 \).

Reactmine uses four hyperparameters which are optimized by grid search to minimize the quadratic loss criterion of Equation (19). SINDy uses one hyperparameter, the sparsity-enforcing penalty coefficient \( \lambda \). It turns out that in all the examples below, there is no value of \( \lambda \) leading to both a good fit and a sparse model. In particular, there is no value of \( \lambda \) for which the hidden dynamics are recovered, as illustrated by Figure S2. For the sake of comparison to SINDy, we thus report the ODE system obtained using the (greatest) value of \( \lambda \) that gives the same quadratic loss as Reactmine.

On the chain CRN example, Table I shows that Reactmine succeeds in recovering the hidden CRN by inferring \( D \xrightarrow{1} E \) first, with the end of the trace as support, as shown in Figure 1. Then the other reactions are learned in backward order, after successive velocity matrix updates, shown in Figure S1 for subtracting the effects of each learned reaction. More details are given in Table S1, where we see that, in this example, the reactions are immediately inferred with the right kinetics and not changed by global re-optimization. Table S2 gives the hyperparameter settings used for the results reported in this section, the number of CRN candidates computed using the best hyperparameter setting found (here 128 chain CRN candidates) the learning time (here 0.31 seconds) and the hyperparameter grid search computation time (here 50 minutes).

In this example, SINDy correctly infers the (ODE terms of the) hidden reaction \( A \xrightarrow{1} B \), then a part of reaction \( B \xrightarrow{1} C \) is present: \( B \) includes the term \(-1.00B\), but only \( 0.18B \) can be found in \( C \) among several other terms that do not correspond to reactions. Likewise, the production of \( E \) by the reaction \( D \xrightarrow{1} E \) is correctly inferred, but the ODE associated with \( D \) is \(-0.37DE\) instead of \(-1.00D\). Moreover, the learned ODEs are not able to generalize the dynamics of species \( D \) and \( E \) on traces that were not used during training, as shown in Figure S3. It is worth noting in this respect that the chain CRN is composed of 8 ODE terms, whereas the SINDy library comprises \( 5 \times (1 + \frac{\lambda}{2}) = 80 \) terms in this example.

The second example concerns the MAPK signaling network, a ubiquitous CRN structure that is present in all eukaryote cells and in several copies. We consider the simplified two-stage (instead of three) CRN model composed of 7 species and 7 reactions of (Qiao et al., 2007). The input species \( A \) goes through a first stage of complexation and phosphorylation to produce the
Table 1: Results obtained by Reactmine and SINDy on the Chain and MAPK CRNs

Using a single simulation trace from one initial state containing the molecular species indicated in bold in the first column. The reactions learned by Reactmine are indicated in green if they belong to the hidden CRN structure, in yellow if they correspond to the kinetics of some hidden reactions, regardless of the precise kinetic constant value as long as the sign is exact, and in red otherwise. For the ODE systems inferred by SINDy on the Chain example, the terms are coloured in green if they correspond to the kinetics of some hidden reactions, regardless of the sign, as shown in Table S1. The dashed red line represents the actual number of nonzero terms in the ground truth CRN. The left arrow shows that the ODE model inferred by SINDy matching the Reactmine loss contains too many terms, around 90, while the right arrow demonstrates that the sparse ODE systems inferred by SINDy are not able to fit the observed velocities.

As shown in Table 1, Reactmine recovers 6 out of 7 reactions of the hidden CRN and 3 other reactions: $ApB \xrightarrow{150} Ap + Bpp$. Signal amplification is caused by the difference of concentrations by several orders of magnitude between the input and the output, as shown in Figure S4. For this example, we set trace parameters $T = 100$ and $\Delta t = \frac{1}{150}$.

On the other hand, the ODE system inferred by SINDy contains two zero-valued differential functions for yet evolving species, and fails on sparsity with

phosphorylated form $Ap$ which plays the role of a kinase on $B$ at the second stage to produce the doubly phosphorylated output species $Bpp$. Signal amplification is caused by the difference of concentrations by several orders of magnitude between the input and the output, as shown in Figure S4. For this example, we set trace parameters $T = 100$ and $\Delta t = \frac{1}{150}$. As shown in Table 1, Reactmine recovers 6 out of 7 reactions of the hidden CRN and 3 other reactions: $ApB \xrightarrow{150} Ap + Bpp$. Signal amplification is caused by the difference of concentrations by several orders of magnitude between the input and the output, as shown in Figure S4. For this example, we set trace parameters $T = 100$ and $\Delta t = \frac{1}{150}$. As shown in Table 1, Reactmine recovers 6 out of 7 reactions of the hidden CRN and 3 other reactions: $ApB \xrightarrow{150} Ap + Bpp$. Signal amplification is caused by the difference of concentrations by several orders of magnitude between the input and the output, as shown in Figure S4. For this example, we set trace parameters $T = 100$ and $\Delta t = \frac{1}{150}$.
an average number of 15 terms per non-zero ODE. Furthermore, the bottom right Figure in Table S1 shows that SINDy fails on this example for all values of its hyperparameter $\lambda$. It is worth remarking in that Figure that the loss is computed by an estimation of the velocity matrix by finite differences on a trace, produced by numerical integration with a more elaborate implicit method, and outflanked by SINDy with low values of $\lambda$. It is worth noting that the 7 ODEs of the MAPK CRN comprise a total of 20 terms, whereas the SINDy library comprises $7 \times (1 + \frac{7 \times 2}{2}) = 203$ terms. Likewise, for the chain CRN, this creates particularly challenging sparse regression problems in the absence of strong independence properties between predictors (Zhao and Yu, 2006). The better results reported in (Mangan et al., 2016) may be explained by the recourse to multiple traces with different zeroes in the initial conditions, similarly to what has been shown in the context of Boolean models in (Carcano et al., 2017).

Table S1 summarizes the results obtained by Reactmine and SINDy on a benchmark of even smaller size CRNs presenting different kinds of difficulties. The learning velocity traces used in those examples, also including the chain CRN, are detailed in Figures S6–S9. The loop CRN adds a feedback reaction $E \rightarrow A$ to the chain CRN, leading to the stabilization of all molecular species on some common concentration value. Reactmine succeeds in recovering the hidden reactions in forward order, directly with the right kinetics. Here again, SINDy recovers some terms of the two first reactions and of the last reaction, but among many other overfitting terms which do not generalize to simulation traces obtained from different initial states, as shown in Figure S10.

The reactant-parallel CRN is just composed of one catalytic reaction, $A + C \rightarrow B + C$, where the catalyst $C$ is produced by two concurrent reactions $D \rightarrow C$ and $E \rightarrow C$. Reactmine first infers the preponderant production of $C$ by $D$, then by $E$, after what the reaction catalyzed by $C$ is correctly inferred. One can notice that the rate constant first inferred for the reaction $E \rightarrow C$ has a small value below its final value by two orders of magnitude. The reason is that right after the inference of $D \rightarrow C$, reaction $E \rightarrow C$ is inferred prior to $E \rightarrow C$. The global re-optimization of rate constants has for effect in this case to set to 0 the rate constant of the second reaction, in favor of the third reaction $E \rightarrow C$ which is thus finally recovered with the right kinetics. In this example, SINDy infers a wrong ODE system that does not reproduce the learning trace, even by increasing the number of time points as shown in Figure S11.

The product-parallel CRN is the symmetrical case of two concurrent consumptions of the catalyst $C$ with the production of species $D$ and $E$, on which SINDy similarly fails. Reactmine first infers the preponderant transformation of $C$ in $E$, then of $C$ in $D$ and then the correct catalysed reaction with little correction of the rate constants by the global optimization phase, as shown in Table S1.

### 3.2 Hyperparameter sensitivity analyses

In this section, we study the impact on the previous results of the hyperparameter values of Reactmine and of the number of time points. Figure S12 shows that Reactmine results are sensitive to both $\alpha$ and $\beta$. In particular, there exist a region for which sufficiently many candidates are proposed and accepted due to a sufficiently high acceptance threshold (e.g. $\beta > 6, \alpha > 0.01$ for the loop CRN). Next, Figure S13 also assesses the sensitivity with respect to $\gamma$ the maximal CRN size and $\delta_{\text{max}}$ the maximum absolute fold change between species variations in a reaction. Only the value of either $\delta_{\text{max}}$ or $\gamma$ is changed, the other hyperparameters are set to values leading to maximum $F_1$-score. $\gamma$ is a sensitive hyperparameter either in terms of quadratic loss value or $F_1$-score. This is expected as being the maximum CRN size, $\gamma$ can be seen as the maximum number of freedom degrees. One should remark that the $\gamma$ value associated with a perfect $F_1$-score is sometimes higher than the size of the hidden CRN. As previously mentioned in Section 3.1 for the case of the Reactant-Parallel CRN, this is due to the inference of reactions whose effect is later set to 0 upon global re-optimization. For this example, these steps were somewhat needed otherwise the ground truth CRN would have been recovered with $\gamma = 3$. It is worth noticing that the hyperparameter sets yielding the lowest quadratic loss are associated with highest $F_1$-score for all CRNs, thus providing empirical evidence concerning the relevance.
of hyperparameter selection based on quadratic loss minimization.

Now, the sensitivity to the number of time points in the trace is evaluated in Figure S14 on the chain CRN. We observe an almost monotonic transition to high F1-score / low reconstruction error as the number of time points increases with optimal performance being reached above 40 time points only.

On the other hand, Figure S11 shows that the failure of SINDy to recover the hidden CRNs persists independently of the number of time points for both the reactant and product-parallel examples. For these examples, increasing up to 200000 time points did not lead to perfect inference as can be seen from the $F_1$-score being different from 1. The learned models display high precision but low recall, suggesting sparse but yet incomplete dynamics. However, the chain and loop CRN could be recovered using 10000 and 500 time points, respectively. This suggests that the problem of sparse regression in that approach rather comes from the high level of correlations observed in single CRN traces without the possibility to vary the zeroes in the initial conditions.

### 3.3 Evaluation on videomicroscopy data

In this section we apply Reactmine to time lapse videomicroscopy data of fluorescent reporters of the cell cycle and circadian clock in NIH-3T3 embryonic mouse fibroblasts (Feillet et al., 2014). These data have been used to develop a coupled model of the cell cycle and the circadian clock in this cell line in Traynard et al. (2016). The cell line was modified to include three fluorescent markers of the circadian clock and the cell cycle: the RevErb-α::Venus clock gene reporter (Nagoshi et al., 2004) for measuring the expression of the circadian protein RevErbα, and the Fluorescence Ubiquitination Cell Cycle Indicators (FUCCI), Cdt1 and Geminin, two cell cycle proteins which accumulate during the G1 and S/G2/M phases respectively, for measuring the cell cycle phases (Sakaue-Sawano et al., 2008). The cells were left to proliferate in vitro in standard culture medium supplemented with 20% of Fetal Bovine Serum. Fluorescence recording was performed in constant conditions with one image taken every 15 to 30 minutes during 72 hours.

A dataset of 67 tracked cells was built from these experiments (Feillet et al., 2014). Figure S15 displays the fluorescence levels trajectories obtained for 3 of these cells as an illustration of the high inter-cell variability and noise level displayed by the data. For this reason, our learning protocol will

- first smooth the curves with a sliding window of 2.5 h;
- apply Reactmine to infer a CRN for each cell individually, using Michaelis-Menten rate functions (as described in Section 2.3.2) in order to fit indirect effects, $\gamma = 3$ in order to discover the main influences between the 3 variables, and the remaining three hyper-parameters selected by grid search as mentioned in Section 2.3.6;
- then compute statistics on the number of reaction occurrences across the $C = 67$ inferred CRNs (one per cell). To this end, we define the mean effect $\mu$ of a reaction $r = (R, P, f)$ on the velocity matrix as:

$$
\mu_r = \frac{1}{nC} \sum_{c=1}^{C} \sum_{l=1}^{n} f(y_l^{(c)})
$$

where $y_l^{(c)}$ is the fluorescence vector at time $t_l$ for cell $c$.

As shown in Figure S16, two reactions clearly stand out compared to the others in terms of effect. The first one, $G2 \rightarrow REV$, represents a possible effect of the cell cycle on the circadian clock through the activation of Rev-Erbo during the G2 phase, in agreement with the main surprising findings of the modeling study of Traynard et al. (2016), using the same dataset. The second most impactful reaction effect-wise is $G1 \rightarrow G2$, the reaction accounting for the cell phase transition from G1 into S/G2/M. The reverse formal reaction $G1 \rightarrow G2$ which could perhaps be expected, is learned but ranked effect-wise behind several meaningless reactions. In terms of occurrence number, one can observe the predominance of a meaningless reaction $G2 + G1 \rightarrow \emptyset$, present in 37 out of 67 cells, yet never inferred first, and with low rate constants.
3.4 Detection of systemic controls on clock gene transcription

In this section, we apply Reactmine to biological data measuring the circadian rhythms of five systemic regulators in mice: body temperature, rest-activity rhythms, food intake, plasma corticosterone and melatonin, as well as the circadian mRNA expression data in the mouse liver of two core clock genes: Bmal1 and Per2. These datasets have been analyzed in [Martinelli et al., 2021] to infer the preponderant systemic regulators on clock gene transcription through another model learning approach. Using an ODE-based model of the liver circadian clock [Hesse et al., 2021] as well as data from four mouse classes, transcription activation profiles $y$ were derived for the two genes. These profiles were regressed on systemic regulators with the aim to infer the significant drivers of clock gene transcription. Let us now attempt to use Reactmine for this task and compare the results.

Since we are looking for an influence model rather than a reaction model properly speaking, we shall use the classical encoding of an influence by one formal reaction catalyzed by the source of the influence [Pages et al., 2018]. We will thus enforce in Reactmine the search of influence reactions of the form $z \to z+y$ where $z$ is one systemic regulator, $y$ one transcription activation profile for a target gene. We shall assume mass action law kinetics representing influence forces.

Furthermore, it is assumed that a systemic regulator $z$ acts either directly, or indirectly though intermediate species by considering its integral counterpart $\int z$, but not both ways. The 5 systemic regulators thus lead to 10 possible influences on Bmal1 and Per2 genes. The Reactmine hyperparameters $\gamma$ and $\beta$ can be set accordingly to 5 and 10. $\delta_{\text{max}}$ is set to 3 as in the previous experiments. Only $\alpha$, the acceptance threshold for influence skeletons, needs to be searched in order to restrict ourselves to preponderant influences only. Figure S17 shows the mean quadratic loss and the number of influences of the CRNs inferred by Reactmine for different values of $\alpha$. We observe that for $\alpha = 6$, the inferred CRNs with only two influences in average reach an MSE below 0.15, i.e. explain more than 85% of the variance.

Remarkably, Reactmine discovers that Food Intake and Temperature are the main influencing factors of clock gene expression, either in a direct or an indirect manner, in agreement with previous findings in [Martinelli et al., 2021]. More precisely, Figure S18 recapitulates the mean kinetic rate constants of the inferred reactions obtained across the traces as a function of $\alpha$. The zoomed part of the figure corresponds to the region where $\alpha = 6$ for which the inferred CRNs contain 2 reactions in average. Using $\alpha = 6$, for Bmal1, the indirect action of Food Intake is deemed the most relevant regulator, followed by the indirect action of Temperature and the direct action of Corticosterone, in agreement with [Martinelli et al., 2021]. Concerning Per2, Food Intake and Temperature again stand out as most important systemic drivers, in both direct and indirect forms. The recovered regulation importance ordering is the same as in [Martinelli et al., 2021] (Figure 6B-C), except for one permutation between $\int \text{Corticosterone}$ and $\int \text{Temperature}$.

4 Conclusion

We have presented Reactmine, an algorithm designed to infer biochemical reactions with kinetics, between molecular species observed in wild type time series data, i.e. without gene knockout or other possibilities to put initial conditions to 0 at will. On a benchmark of hidden CRNs of increasing difficulty, including one model of MAPK signal transduction [Qiao et al., 2007], we have shown that Reactmine is able to recover the hidden CRN, or an essentially equivalent form of it, from one single ODE simulation trace. On the opposite, the state-of-the-art sparse regression algorithm for non linear dynamical systems, SINDy [Brunton et al., 2016], with appropriate function libraries for the examples, fails to infer sparse ODE systems from such wild type traces, and even to reproduce other simulation traces obtained from different initial states.

The behavior of sparse regression algorithms is indeed conditional to assumptions about the low level of correlation between predictors [Zhao and Yu, 2006]. Those hypotheses are not satisfied in the context of wild type time series data, specifically in a low data regime. The possibility of varying the traces by setting to 0 some initial conditions has a de-correlation effect which may explain the better results reported in [Mangan et al., 2016], similarly to what has
been shown for boolean models in [Carcano et al. (2017)]. More work is however needed to develop those arguments in the general context of sparse identification of non linear dynamics.

Reactmine solves those issues by not relying on sparse regression but on a bounded-depth search of reaction candidates, inferred in a sequential manner on a subset of supporting transitions, with statistical criteria about the inferred velocities. This leads to four hyperparameters which are currently chosen by grid search for the three sensitive ones. Some formulae are currently under investigation for providing them with default values as a function of the dimension of the problem. The restriction to 0/1 stoichiometry adopted in this article can be easily dropped as long as a limited number of reaction schemas is considered. When applied to real biological data of videomicroscopy data and systemic circadian controls of clock genes, we have shown that Reactmine is able to retrieve the main conclusions of the model-based analyses done respectively in [Traynard et al. (2016)] and [Martinelli et al. (2021)] on the same datasets.

These encouraging results should motivate applying Reactmine to new study cases on the one hand, and on the other hand, investigating extensions of our approach to deal, in that setting, with the open problem of learning models including latent species.

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Supplementary material

S1 Detailed results of Reactmine and SINDy

Computation times reported here have been obtained on a Macbook M1 2020 13” with 8 cores. For Reactmine, the grid search was parallelized. For [SINDy](https://pysindy.readthedocs.io) we used the [pysindy library](https://pysindy.readthedocs.io) with STLSQ (sequential least square thresholding) optimizer, as we observed that it yielded better results than Lasso or SR3. We chose to report in Table S1 and in the main text, the ODE system inferred by SINDy with a value of regularization hyperparameter $\lambda$ that leads to the quadratic loss (which is not zero due to numerical integration errors) of the ground truth CRN recovered by Reactmine.
| Name | Hidden CRN | CRN inferred by Reactmine | ODE system inferred by SINDy |
|------|------------|----------------------------|------------------------------|
| Chain | A \xrightarrow{1} B | D = 0.9936 \quad E = 0.9941 | \( A = -1.064 \) |
| Loop | A \xrightarrow{1} B | C = 0.9936 \quad D = 0.9974 | \( B = 1.014 \quad D = -0.06AB \) |
| | C \xrightarrow{1} D | B = 1.0904 \quad E = 0.9952 | \( C = 1.06 \quad D = -0.06\) |
| | D \xrightarrow{1} E | A = 1.0000 \quad B = 1.0000 | \( D = -0.06\) |
| | A \xrightarrow{1} B | C = 0.9936 \quad D = 0.9974 | \( A = -1.064 \) |
| Reactant Parallel | A \xrightarrow{1} C \xrightarrow{1} B + C | D = 0.9936 \quad E = 0.9974 | \( A = -1.124 \quad 5100000276790.22C = 0.87\) |
| | D \xrightarrow{1} C | B = 0.9936 \quad E = 0.9974 | \( B = -1.064 \) |
| | E \xrightarrow{1} C | A = 0.9936 \quad B = 0.9974 | \( C = 0.9936 \) |
| Product Parallel | A \xrightarrow{1} C \xrightarrow{1} B + C | D = 0.9936 \quad E = 0.9974 | \( A = -0.06\) |

**Table S1:** Results obtained by Reactmine and SINDy on hidden CRNs using a single simulation trace from one initial state containing the molecular species indicated in bold in the first column. The learned reactions are indicated in green if they belong to the hidden CRN, in yellow if they lead to equivalent terms of the associated ODEs, and in red otherwise. For a learned reaction, the number written underneath the arrow is the initial rate constant value learned with the reaction, before global re-optimization. For the ODE systems inferred by SINDy, the terms are coloured in green if they correspond to the kinetics of some hidden reactions, regardless of the precise kinetic constant value as long as the sign is exact, and in red otherwise.
Table S2: Reactmine computation times and hyperparameter values used for the results reported in Table S1. The best hyperparameter settings indicated in the second column for each example, are found by grid search with a range of values reported in the third column, and a computation time given in the last column. The CRN learning computation times in these settings are given in the third column, with the number of generated CRN candidates in the fourth column.
Figure S1: Successive velocity trace updates done by Reactmine for recovering the chain CRN.
Figure S2: Quadratic training loss (blue) and number of nonzero terms in the library (red) found by SINDy as a function of $\lambda$ in our benchmark of reactant-parallel, product-parallel, chain, loop and MAPK CRNs. The dashed red line represents the actual number of nonzero terms in the ground truth ODE associated to each hidden CRN. The dashed darkblue line stands for the (non-zero due to numerical errors) quadratic loss value of the ground truth CRN.

Figure S3: Simulation of the ODE model learned by SINDy in the Chain example using different initial conditions (dotted line) compared with ground truth (solid line), showing erroneous dynamics for $D$ and $E$. 
Figure S4: **Learning trace and velocity trace for the MAPK CRN.** The lower panel shows the same plots with concentrations and velocities normalized by their maximal values along the trace for visualization purposes.
Figure S5: Reproduction of the MAPK CRN simulation traces obtained from different initial conditions using the CRN inferred by Reactmine. Time-resolved species concentrations are normalized to their maximal value across the trace. $y_0^{(1)} = (2, 0, 0, 0, 0, 0, 0.5, 0.01)$; $y_0^{(2)} = (0, 0, 0, 0, 0, 0, 0, 0.1)$; $y_0^{(3)} = (0, 0, 0, 0, 0, 0, 1.5, 0.001)$. The second column displays the loss between the simulation traces of the MAPK CRN and the CRN inferred by Reactmine.
Figure S6: Learning trace and velocity trace for the Reactant-Parallel CRN.

Figure S7: Learning trace and velocity trace for the Product-Parallel CRN.

Figure S8: Learning trace and velocity trace for the Chain CRN.
Figure S9: Learning trace and velocity trace for the Loop CRN.

Figure S10: Simulation of the ODE model learned by SINDy in the Loop example using different initial conditions (dotted line) compared with ground truth (solid line), showing erroneous dynamics on all species.
Figure S11: Sensitivity of SINDy to the number of time points with $\lambda$ selected to yield the highest binary $F_1$-Score (i.e. $F_1$-score with labels 1 for terms with non-zero coefficient in the ODEs, independently of their sign, and labels 0 for zero coefficient). Precision, recall, F1-score and quadratic loss are displayed as a function of the number of time points in the learning simulation trace used. With very high number of time points, SINDy succeeds in recovering the hidden chain and loop CRNs, but this is not the case for the reactant-parallel and product-parallel CRNs.
Figure S12: Sensitivity of Reactmine to $\alpha$ and $\beta$ hyperparameters. Quadratic loss $\|\hat{V} - F(Y, k)\|_F^2$ (lower is better) and F1-score (in parenthesis, higher is better). The colorbar levels relate to the quadratic loss. $\delta_{\text{max}} = 3$ and $\gamma = 6$ except for the reactant-parallel CRN where $\gamma = 5$, as it yielded a better result.
Figure S13: Sensitivity of Reactmine to $\delta_{\text{max}}$ and $\gamma$ hyperparameters. The quadratic loss is reported in darkblue, the $F_1$-score in red, negated for visualization purposes.
Figure S14: Sensitivity of Reactmine to the number of time points for the chain CRN with $\delta_{\text{max}} = 3, \alpha = 0.2, \beta = 7, \gamma = 5$. The time horizon of the simulation is $T = 10$. The quadratic loss is reported in dark blue, the $F_1$-score in red. The x-axis is in log scale.
Figure S15: Fluorescence videomicroscopy plots of 3 cells among the 67 cells of the videomicroscopy data. Traces have been smoothed with a moving average.
Figure S16: **Results of Reactmine on videomicroscopy dataset.** Bars in green report the mean effect of each reaction across time and cells in the videomicroscopy data. Standard deviation is indicated across the cells. The two reactions with the most important effect are in accordance with the previous findings done in studies of this dataset. The number of occurrences of a reaction is plotted in red.

Figure S17: **Results of Reactmine on systemic regulators of clock gene expression.** Average mean square error (green) and number of reactions (red) for the CRNs inferred by Reactmine are given as a function of $\alpha$. The statistics are computed over the $N = 2000$ trajectories $\{y^{(j)}\}_{1 \leq j \leq N}$ considered. The green dotted horizontal line represents a MSE threshold of 0.15, equivalent here to 15% left unexplained by the inferred CRN. The red dotted vertical line reveals that for both *Bmal1* and *Per2*, CRNs with only two reactions are enough to reach a MSE below 0.15.
Figure S18: Systemic drivers of clock gene transcription inferred by Reactmine. The mean rate constant associated to a catalyzed reaction is reported as a function of $\alpha$. The mean is computed over the $N = 2000$ trajectories $\{y^{(j)}\}_{1 \leq j \leq N}$. The plot zoom corresponds to a region close to $\alpha = 6$. 

\[\text{Kinetic rate } \hat{k} \text{ associated with catalyzed reaction}\]