Dynamical differential expression (DyDE) reveals the period control mechanisms of the Arabidopsis circadian oscillator

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Abstract

The circadian oscillator, an internal time-keeping device found in most organisms, enables timely regulation of daily biological activities by maintaining synchrony with the external environment. The mechanistic basis underlying the adjustment of circadian rhythms to changing external conditions, however, has yet to be clearly elucidated. We explored the mechanism of action of nicotinamide in Arabidopsis thaliana, a metabolite that lengthens the period of circadian rhythms, to uncover key transcriptional mechanisms of circadian period. To identify the key mechanisms involved in the circadian response to nicotinamide, we developed a systematic and practical modeling framework based on the identification and comparison of gene regulatory dynamics. Our mathematical predictions, confirmed by experimentation, identified key transcriptional regulatory mechanisms of circadian period and uncovered the role of blue light in the response of the circadian oscillator to nicotinamide. We suggest that our methodology could be adapted to predict mechanisms of drug action in complex biological systems.

Author summary

Circadian oscillators are internal timing devices that produce rhythms with a period of about 24 hours to regulate a wide range of biological processes for nearly all organisms. Circadian oscillators adjust phase and period in response to external cues such as light and temperatures and internal cues such as metabolites to maintain synchrony with the daily timing of light and dark cycles resulting from Earth’s rotation. We investigated the dynamical response of the Arabidopsis thaliana circadian network to nicotinamide, a metabolite that lengthens the period of circadian rhythms, to uncover key transcriptional
mechanisms required for the adjustment of daily biological activity. The identification of the mechanisms of drug response is challenging, as a complex cascading effect causes large section of the transcriptome to be differentially expressed, despite not being directly affected by the drug. To identify the source of the change in circadian period, we introduce a modelling strategy based on the identification and comparison of gene regulatory dynamics before and after the perturbation occurred. The Dynamical Differential Expression (DyDE) methodology uses a reverse engineering approach that favours both the identification of unknown Gene Regulatory Network (GRN) topology and the interpretation of the possible dynamical changes, without the need to cover extensive experiments or to make prior assumptions of network dynamics. Subsequently, we show that our methodology can reliably identify the source of a perturbation in complex regulatory systems such as the circadian network. The proposed mathematical framework is scalable and flexible, so that it can be applied to large datasets with scarce sampling.

Introduction

The synchronization of physiological rhythms with the external environment is important for nearly all organisms. Circadian oscillators are internal timing devices that produce rhythms with a period of about 24 hours to regulate a wide range of biological processes. Circadian rhythms maintain synchrony with the daily timing of light and dark cycles resulting from Earth’s rotation by constantly integrating environmental signals. This process of synchronization is called entrainment. Studying the mechanisms that dynamically adjust circadian period and phase, therefore, is critical to understand the control of daily biological activities.

In Arabidopsis thaliana, the circadian oscillator consists of a complex circuit of highly connected transcriptional regulators. Together, they coordinate global transcript accumulation and diverse biological processes, such as photosynthesis, hormone signaling, hypocotyl elongation and plant-pathogen interactions [1,2,3,4,5]. The light perception of the circadian oscillator is conferred by a suite of photoreceptors. The photoreceptors are split into two classes: phytochromes (principally PHYA and PHYB), that primarily sense the red portion of the spectrum [6] and cryptochromes (CRY1 and CRY2) that are sensitive to blue light [7,8,9].

Recent studies have demonstrated a role for metabolism in regulating and entraining the circadian oscillator of Arabidopsis thaliana. The primary metabolite sucrose accelerates the circadian oscillator (i.e., reduces its period) through regulation of the morning expressed gene PSEUDO RESPONSE REGULATOR (Prr) [10], while GIGANTEA (GI) has been identified as a necessary sucrose-signaling mediator in the dark [11]. Another metabolite, nicotinamide (NAM), a breakdown product of nicotinamide adenine dinucleotide (NAD), causes long period of the circadian oscillator in all organisms tested [12,13]. The mode of action of NAM is uncertain: various mechanisms having been proposed, including inhibition of the production of the Ca²⁺-agonist cyclic adenosine diphosphate ribose (cADPR), inhibition of polyADP ribose polymerases and histone modifications [12,13,14]. The goal of this study was to use NAM as a tool to identify the processes responsible for a change in circadian period, which might be required for circadian entrainment and homeostatic adjustment [15,16,17].

The discovery of drug modes of action, however, is still a costly and inefficient process, which often requires considerable prior knowledge of a biological system and/or a vast amount of data in several experimental condition (e.g. mutations). A major difficulty is the complex ripple effect of treatments affecting transcriptional networks. Large sections of the transcriptome can be differentially expressed, despite not being directly affected by the treatment (off-
targets) (Fig 1). Due to the large number of feedback loops involved in a complex and relatively small Gene Regulatory Network (GRN), such as the circadian clock, this effect is particularly significant as a perturbation anywhere in the network typically strongly affects all molecular concentrations. Additionally, as the perturbations induced by NAM in the circadian clock are intrinsically related to changes in circadian period, a large part of the transcripts will typically be differentially expressed. Thus, Differential Expression (DE) analysis, the traditional approach used to identify the mechanisms that alter biological behavior in response to drugs, environmental signals or genetic lesions [20], will usually fail to identify the small number of genes central to the biological perturbation. The main reason is that DE only performs statistical analysis of changes in gene expression levels [21,22]. As an alternative to the DE analysis, we devised a modeling framework that identifies and characterizes differentiated regulatory dynamics between genes to capture key mechanisms involved in NAM-induced perturbations in the circadian system of Arabidopsis. The rationale behind this approach is that not only genes, but also their interactions, are affected by a drug. This reasoning is further supported by [18,23,24], which highlight the fact that drugs and diseases mechanisms should be regarded as network instead of gene-centric perturbations. We designed our modelling strategy so that it could be applied to scarce data without the need to cover extensive experiments or to make prior assumptions of network dynamics. In particular, we consider only gene expression data with and without NAM.

Fig 1. Treatment effects in transcriptional networks. Treatment effects can be perceived as perturbations in molecular networks [18,19]. In transcriptional networks, such perturbations usually only affect a very small number of regulatory links directly. For example, only the red links have been directly affected by the treatment. All other links are unchanged, although all nodes (concentrations) in the Figure have been (indirectly) affected due to cascading and feedback effects. Hence, Differential Expression (DE) might not distinguish between direct and indirect effects of a treatment. Dynamical Differential Expression (DyDE), therefore, investigates how and why changes occur, instead of simply measuring what and how much is produced by those changes.

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On one hand, complex nonlinear models have the potential to capture the dynamical relationships between genes with great precision. A successful application of Michaelis-Menten dynamics to reverse engineer network topology from multiple experiments, circadian time-series data is presented and compared to state-of-the-art strategies in [25]. Alternatively, a community-driven comparison of (non)linear approaches (e.g., mutual information-based, Bayesian networks, random forests, neural-networks, etc.) for the inference of (non-circadian) gene regulatory networks has been achieved in [26,27]. On the other hand, high model complexity can lead to overfitting (fitting the noise instead of the dynamics) without sufficient data or detailed knowledge such as network topology, types of nonlinear interactions, or potentially some of the model parameters (e.g., Hill coefficients). As for non-model based methods, such as [28,29,30,31], it is not clear how they can be used to compare subtle changes in dynamics caused by perturbations, and pinpoint the source of those perturbation. We developed, therefore, a systematic and scalable dynamical modeling framework named Dynamical Differential Expression (DyDE). DyDE uses a black box-type modeling approach to reverse-engineer simple yet consistent and comparable gene regulatory dynamics from time-series data. In addition, it does not use any prior information and, hence, it is unbiased towards prior knowledge of network topology and dynamics. By comparing changes in both topology and subtle dynamic modifications of regulatory mechanisms, we were able to considerably narrow down potential targets of NAM in the circadian clock.

The findings predicted by DyDE are experimentally tested and demonstrate the role of the circadian gene PRR7 as a key regulator of dynamics adjustment of the circadian clock. In addition, TIMING OF CAB EXPRESSION 1 (TOC1) and the interplay between PRR7 and PSEUDO RESPONSE REGULATOR 9 (PRR9) are identified as the main mediators of the circadian system response to NAM.

The modeling insights also identified alterations in CRY2 dynamics resulting from the NAM treatment. Therefore, we also investigated the role of blue light in the circadian period change of NAM-treated plants. In particular, we found that blue light regulates circadian oscillations of [Ca^{2+}]_{cyt} through a NAM-sensitive pathway. These new perspectives contribute to the understanding of the mechanistic details underlying the regulation of period of circadian oscillators.

Overall, the results suggest that DyDE is a useful tool to generate reliable hypothesis from time-series data for the identification of drug targets in complex biological systems.

**Methods**

To investigate how NAM might regulate the period of the circadian oscillator we first used statistical tools to identify those transcripts that have circadian rhythms in abundance in both untreated and NAM-treated plants. Then, we introduce the Dynamical Differential Expression (DyDE) approach to characterize altered dynamics within the circadian regulatory network of NAM-treated plants.

The hypothesis generated by DyDE were experimentally tested using genetic mutant and physiological experiments in different light conditions. Finally, we extended DyDE to the whole rhythmic transcriptome to further investigate clock period regulation.

**Statistical characterization of circadian transcripts**

To assess whether genes are regulated by the circadian oscillator, most methods take advantage that circadian regulation of transcript abundance resemble a sinusoid. To estimate circadian period of the regulation of a particular transcript, the main idea is to find the sinusoid that most closely matches its abundance over time [32,33]. However, in NAM-treated plants the
changes in abundance of circadian-regulated transcripts have a considerable number of non-sinusoidal profiles (S1 Fig). To overcome this problem, we devised a learning approach based on pseudo-sinusoidal functions to properly assess the rhythmicity and the corresponding circadian period of signals from gcRMA normalized microarray data of NAM treated plants. To infer period, phase and amplitude, linear trends are eliminated by removing the best straight-line fit and pseudo-sinusoidal functions are fitted to each signal to minimize the 2-norm error. Pseudo-sinusoidal functions account for many signals that are periodic but not sinusoidal. Pseudo-sinusoidal functions are constructed by joining together two sinusoids with different periods. Hence, a complete oscillation of a pseudo-sinusoidal function consists of the first sinusoid (of period \( p_1 \)) in the first half-oscillation, and the second sinusoid (of period \( p_2 \)) in the second half-oscillation (Fig 2A). The resulting period of the pseudo-sinusoidal function is defined as \( p = \frac{p_1 + p_2}{2} \).

\[
S = \begin{cases} 
A \sin \left( \frac{2\pi}{p_1} t + \varphi_1 \right), & t \in \left[ 0, \frac{p_1}{2} \right] \\
A \sin \left( \frac{2\pi}{p_2} \left( t - \frac{p_1}{2} + \frac{p_2}{2} \right) + \varphi_1 \right), & t \in \left[ \frac{p_1}{2}, \frac{p_1}{2} + \frac{p_2}{2} \right]
\end{cases}
\]

where \( A \) is a scaling factor that accounts for the amplitude of the signal and \( \varphi_1 \) is the phase of the signal. The algorithm searches possible combinations of \( p_1 \) and \( p_2 \) to minimize the least square distance between pseudo-sinusoidal functions and the data. We allowed periods \( p_1 \) and \( p_2 \) to vary between 12 and 36 hours. A perfect sinusoid gave a high fit for the wild-type background dataset. We found that three periodic signals were highly represented in the dataset. In particular, those with \( p_1, p_2 \) equal to: \( \frac{p}{2}, \frac{p}{2} \) (pure sinusoid); \( \frac{p}{2} + 3.8, \frac{p}{2} - 3.8 \) (\( p_1 \) is greater than \( p_2 \)); and \( \frac{p}{2} - 7.3, \frac{p}{2} + 7.3 \) (\( p_1 \) is smaller than \( p_2 \)) (Fig 2A).

We used a logistic regression framework to generate a probabilistic discriminative model that estimates the probability of a gene to be rhythmic given its time course data. In this case, the classification problem only contains two classes: rhythmic (\( C_1 \)) and arrhythmic (\( C_2 \)). For each transcript, a set of 8 features \( x = \{ X_1, X_2, \ldots, X_8 \} \) is computed and empirically believed to be crucial to distinguish between rhythmic and arrhythmic transcripts.

The features were computed from 2 signals: the first signal (A) corresponds to the average of replicates and (B) being a single replicate for which the L2-norm error with the best fitted pseudo-sinusoidal function is lower than for the other replicate. The following features were computed: ratio of power in the 18–32 hours frequency range (of (A) and (B)), L2-norm of the error to the best fit of pseudo sinusoidal function (of (A) and (B)), the variance of the power spectrum (of (A) and (B)) and the amplitude of the best fitted pseudo-sinusoidal function (of (A) and (B)).

The log of the ratio of probabilities between the two classes, also known as the log odds, is given by [34]:

\[
\log \left( \frac{p(\text{rhythmic}|x)}{p(\text{arrhythmic}|x)} \right) = \log \left( \frac{p(C_1|x)}{p(C_2|x)} \right) = \log \left( \frac{\sigma}{1 - \sigma} \right) = \logit(\sigma)
\]

The goal of the logistic regression is to estimate \( \sigma \) for a linear combination of the \( X_n \) features such that:

\[
\logit(\sigma) = b_0 + b_1X_1 + b_2X_2 + \cdots + b_nX_n
\]

The weights \( b_i \) of the independent variables \( X_i \) were estimated using the \textit{mnrfit} function in MATLAB. The algorithm is initially trained with a mix of 100 rhythmic and 100 arrhythmic transcripts randomly chosen from the dataset and visually inspected to show clear (ar)
Finally, the decision boundary was set so that if \( p(C_1|x) > 0.5 \), the gene was classified as rhythmic, and vice versa. Our approach, therefore, is inspired by the patterns observed in the dataset but not strictly constrained to pure cosine shapes. With the inclusion of the \( S \) function, we allow the search for asymmetric signals, which represent a large part of the transcriptome. A main distinction with the previously introduced algorithms, therefore, is the data-specific, learning approach devised to allow for a wider range of periodic signals. However, this offers additional advantages such as a dedicated way to handle noise between replicates, or the information in the frequency domain of the signal, which are both learned from the data. Comparison of performances with standard periodicity assessment tools is shown on S2 Fig.

**Network inference and analysis by DyDE**

Like most biological systems, circadian clocks have a large number of feedback loops. Hence, a perturbation anywhere in the network typically affects all nodes (in this case, their molecular
concentration and time profiles), which makes the problem of inferring the entry point of a perturbation hard using DE analysis. We proposed, instead, that key mechanisms involved in NAM-induced perturbations in the circadian system of Arabidopsis can be captured by identification and comparison of regulatory dynamics before and after the perturbation occurred. Assume that a perturbation, such as NAM, changes the regulatory dynamics between two genes (e.g. by binding to a transcription factor) while leaving intact the rest of the system. Due to feedback interconnections, all the clock genes would change their expression, which, in turn, would change the expression of all circadian genes. While thousands of genes change their expression, only one regulatory link was actually affected. Our goal is to find this link (or links, in case of multiple perturbation entry points). To achieve this, we developed DyDE that looks for changes in links, instead of nodes. DyDE uses Linear Time-Invariant (LTI) models, a type of black box model, to systematically capture the dynamics underlying the biochemical mechanisms of circadian gene regulation, without relying on a priori knowledge of the system or extensive database. Such models benefit from a rich theory and a well-established collection of tools that makes the analysis of its dynamical properties straightforward, as contrast to detailed mechanistic models. In addition, the estimation of the parameters of such models is reliable and computationally efficient. The description of biological mechanisms of the Arabidopsis circadian clock from time-series data by LTI models has been studied in [35]. More recently, the performances of such linear modelling approach to reverse engineer the clock topology were compared for two extensively used Arabidopsis oscillator models [36] and confirmed that the majority of oscillator links can be represented by simple linear dynamics. However, the use of LTI models to detect dynamical perturbation in the gene regulatory network resulting from chemical treatments is novel.

The first step of DyDE consists of uncovering dependencies and quantifying dynamics between genes with LTI models. Our mathematical framework estimates a collection of Single Input-Single Output (SISO) models between pairs of genes to characterize the system dynamics. The limited number of available time points restricted the modelling of SISO systems to first and second order models. Overall, second order systems did not improve significantly the fitness of models and resulted in a considerable increase of false positives (overfitting). Hence, in this analysis of the circadian system, we restrict the model order to one. Mathematically, the dynamics between two genes can be represented as:

\[
\frac{dy(t)}{dt} = a\ u(t) - b\ y(t) + c
\]

where \(u(t)\) and \(y(t)\) represent the time series of the regulatory gene and the regulated gene, respectively. In addition, \(b\ y(t)\) corresponds to the degradation rate of gene \(y\), \(a\ u(t)\) corresponds to the influence of \(u(t)\) on the rate of \(y(t)\) and \(c\) is a constant offset. System identification is performed using the function ‘pem’ implemented in MATLAB to minimize the prediction error [37]. The model has a total of three parameters (\(a\), \(b\), and \(c\)), leading to efficient solutions. We chose a subspace initialization algorithm since it performed similarly as randomizing initial conditions—for the vast majority of models (99%), the final solution was identical with either method. This suggests that the chances of being trapped into a local minimum are negligible.

The estimation of parameters requires low computational time: a single system between a pair of genes is typically identified within few seconds (Intel Core i5). This modeling is independently repeated for all available pairwise genes, where each gene takes its turn as being an input and then an output to another gene. This modeling approach, therefore, generates a large amount of SISO LTI models (\(n^2 - n\) models, where \(n\) corresponds to the amount of genes, and self-regulation is not considered) to describe the system. Each potential link between two
genes is validated if the corresponding model reproduces the dynamics involved with a sufficient degree of precision, which is characterized by a high goodness of fit, defined as:

$$\text{fitness} = 100 * \left(1 - \frac{\sum_{k=1}^{N} \sqrt{(y - \bar{y})^2}}{\sum_{k=1}^{N} \sqrt{(y - \bar{y})^2}}\right)$$

where \(y\) is the validation data, \(\bar{y}\) is the average value of the validation data, and \(\widehat{y}_k\) is the estimated output. MATLAB function \textit{compare} can be used to compute the fitness of the model. A fitness equal to 100% corresponds to a perfect identification. The choice of such metric is motivated by the dependency of noise towards the abundance of gene expression. When the distance of the true data points towards the mean is large (represented by the denominator in the above equation), the fitness conveniently penalizes less the error term, which lies in regions where the intrinsic noise involved in the gene expression is potentially the largest.

The second step consists in identifying the effect of a treatment, NAM in our case, on the biological network. While a treatment might affect the abundance of many transcripts, only a few links are affected, as depicted in red in Fig 1. Hence, checking whether links are affected before and after perturbation can potentially lead to the finding of the entry point of the treatment. For this purpose, two cases are of particular interest. First, a link between two genes is validated in the untreated system alone (i.e. it is not possible to find a combination of \(a\), \(b\) and \(c\) so that the model in the treated system provides a good match with the data anymore). Second, a link is validated in both systems, but the way one gene regulates the other may change; this is a much subtler change in the dynamics of the link. The latter case requires us to compare the dynamics between both links. Here, we use a rigorous and well-established tool from engineering known as the nu-gap [38]. Originally developed to address the stability properties of closed loops systems defined in the same feedback loop, the nu-gap essentially measures the distance, from a perturbation point of view, between linear models. This property is particularly relevant in the context of circadian clock networks, which consist in regulatory networks with several feedback loops. This then facilitates us to determine the significance of the dynamical change of a link between experimental conditions. The nu-gap returns a value between 0 to 1, quantifying whether the models are similar or very different, respectively. [39] have suggested that values above \(\sim 0.2\) could be used to infer the main target of a perturbation. The nu-gap is computed using the \textit{gapmetric} function in MATLAB. It should be applied to all models that have been estimated in both networks. If the signals are concentrated around a particular range of frequencies (such as oscillating signals), the gap should be measured 'locally' around that range of frequencies only, since they dominated the model estimation in Step 1.

Next, we explain the key ideas behind DyDE through a small number of genes in the Arabidopsis circadian oscillator. For example, the following model considers \(TOC1\) as an input and \(PRR9\) as an output.

$$\frac{d[PRR9]}{dt} = a[TOC1]_i - b[PRR9]_i + c$$

where \(b\) represents the strength of activation or repression induced by \(TOC1\) on the expression rate of \(PRR9\), and \(a\) corresponds to the degradation rate of \(PRR9\). These parameters are estimated by minimizing the prediction error from the untreated time-series for both \(TOC1\) and \(PRR9\).

In this case, we found a model in good agreement with the data (57% fitness), suggesting that indeed \(TOC1\) regulates \(PRR9\) (Fig 3A). Moreover, the model demonstrates that the rate of change of the concentration of \(PRR9\) is proportional to the concentration of \(TOC1\). Note that the other way around (i.e., \(PRR9\) regulating \(TOC1\)) could not be established since the
respective model has a low goodness of fit (16%, Fig 3B). These results are consistent with the literature [40]. Hence, we would then establish a link from TOC1 to PRR9, but not the other way around (Fig 3C).

Then, a model is estimated between TOC1 and PRR9 from the NAM-treated time-series. From the untreated and treated time-series alone, it is unclear whether the link dynamics have changed (Fig 3D). The optimal model parameters, however, have significantly changed. A null gap of ~0.5 confirms that indeed the link has been affected. This result indicates that there is large perturbation in the regulatory dynamics that links TOC1 to PRR9, which, therefore, should be considered as a strong candidate for being an entry point for NAM in the system. If true, knocking down either TOC1 or PRR9 would therefore lead to NAM no longer affecting...
the clock. This analysis is then repeated for all common links between untreated and treated plants.

Results
We identified 3859 (18.4%) circadian-regulated transcripts for the untreated plants (Fig 2A). These were enriched for Gorilla terms 'Circadian Rhythm' and 'Rhythmic Process' ($p = 4.07\times10^{-18}$; S2 Table; GEO No. GSE19271). A total of 2588 (12.3%) transcripts were identified as rhythmic in NAM-treated plants (Fig 2A), with a mean increase in period from $24.0 \pm 2.1$ h (-NAM) to $27.4 \pm 2.6$ h (+NAM) (Fig 2B) and without a noticeable change in amplitude (Fig 2C).

DyDE applied to the Arabidopsis circadian clock genes
We considered a total of 17 known clock genes: CCA1, LHY, PRR9, PRR7, PRR5, RVE8, GI, TOC1, ZTL, ELF4, ELF3, PHYA, PHYB, CRY1, CRY2, CHE and PRR3. However, the core oscillator genes ZTL, ELF3, PHYB, CRY1, PRR3 and CHE were identified as non-rhythmic in the presence of NAM, which was confirmed by visual inspection (S1 Fig). Hence, these genes are excluded from the modeling of NAM targets as they cannot be contributing to the rhythmic dynamics of the remaining oscillator components that are measured in the presence of NAM.

As a first step, we computed models for all available pairs of the clock genes for both conditions, totaling 220 SISO models (110 in untreated and 110 in NAM). We kept only those models with good agreement with the data, i.e. above a fitness threshold. On one hand, the user-defined threshold has to be set large enough to reliably capture the dynamics involved between genes, and provide the nu-gap analysis with comparable models. On the other hand, the threshold has to be set sufficiently low to consider enough gene-to-gene relationships to detect a dynamical perturbation in the network. Here, the fitness threshold was set to 46% as we noted that below this threshold, the amount of unknown regulations dramatically raised (S3 Fig; S3 Table).

In total, 70 regulatory links were retained for untreated plants and 55 links for NAM-treated plants between the 11 clock genes. The untreated models describe 70% of the known regulatory pathways among these 11 genes (S3 Table; S3 and S4 Figs [40]). 64% of which, had the expected activation or inhibition effect. These numbers are remarkable, taking into account the model simplicity, and confirms that the majority of clock links can be represented by simple linear dynamics [35,41,42].

In particular, 28 links were present in the untreated samples but not in the NAM-treated samples. These 28 links form a network from now on referred to as "regulation loss" network, which captures the links abolished by NAM. In addition, 42 links are present in both conditions which form a network, so called "common" network that is common to both treated and untreated plants (S3 Table).

We used the nu-gap to identify those links among the common network whose dynamics were significantly affected by NAM. Fig 4A and S4 Table depict the comparison of the dynamics of each link with the nu-gap. All regulatory interactions are somehow affected by the treatment, which is expected from the interconnected circadian network. Let us then consider the highest nu-gap values, which are associated with the following links: TOC1 to PRR9 (0.5), those originating from CRY2 to ELF4 (0.47), LHY (0.42) and RVE8 (0.37) and PRR9 to CRY2 (0.35). Interestingly, the only inferred interaction originating from CRY2 that does not seem affected connects to TOC1 (nu-gap of 0.06). These results suggest that a major dynamical change is induced to CRY2 in the dynamical response of the circadian clock to NAM. In addition, the largest nu-gap value suggests that the causality within the time course data of TOC1
PRR9 has changed significantly differently towards the treatment, as compared to the other parts of the circadian network. We then used a standard network topology metric to identify the genes that are central to the drastic changes in dynamics captured by the regulation loss network. This topology metric

Fig 4. DyDE applied to the Arabidopsis circadian oscillator genes. (A) Common network and nu-gap analysis. The common network displays the models that have been validated in both untreated and NAM-treated plants. A directed arrow from gene a to gene b (blue circles), therefore, represents a dynamical model that captures the dependency of b on a. Red arrows represent the models associated with the top five highest nu-gap values. (B) Bar plot comparing the connectivity loss (%) associated to each gene. For a particular gene, the connectivity loss corresponds to the total amount of incoming and outgoing links that were validated in untreated plants but not in NAM-treated plants. Error bars represent the standard deviation of connectivity loss for ± 5% change in fitness threshold selection.

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and PRR9 has changed significantly differently towards the treatment, as compared to the other parts of the circadian network.

We then used a standard network topology metric to identify the genes that are central to the drastic changes in dynamics captured by the regulation loss network. This topology metric
accounts for the connectivity of a gene, i.e. the number of its incoming and outgoing links. This measure is estimated for each gene of the regulation loss network. As an example, PRR7 has six incoming links and nine outgoing links for untreated plants. The connectivity of PRR7 in untreated plants is then equal to 15. Among those, only six of were present in NAM-treated plants. PRR7, therefore, has a connectivity of nine in the regulation loss network, which correspond to a loss of 60% of its connectivity from untreated to NAM treated plants. As a result, CCA1 (61%), PRR7 (60%), TOC1 (57%) exhibit the highest connectivity drop (Fig 4B; S5 Table). This result identifies the biological functions of CCA1, PRR7 and TOC1 as being highly affected by NAM in the regulation of the circadian clock.

DyDE, therefore, identifies the regulatory dynamics of TOC1-CRY2-CCA1-PRR7 as being predominately affected by NAM as a result of both nu-gap and connectivity analysis. Accordingly, the strong emergence of the blue light receptor CRY2 in the nu-gap analysis suggests that nicotinamide alters the regulation of the interactions between light signaling and the circadian oscillator. These findings are further examined through mutant analysis and single wavelength light experiments.

PRR7/PRR9 inter-regulation together with TOC1 are targets of nicotinamide

To test the predictions that TOC1, CRY2, CCA1 and PRR7 are associated with the effect of NAM on the circadian oscillator, we experimentally investigated the sensitivity of circadian mutants to NAM. All mutants responded to NAM with increased circadian periods, with the exception of two independent lines of the same T-DNA insertion allele of PRR7, which were insensitive (prr7-3 p > 0.95; prr7-11 p > 0.95 Fig 5A and 5B; S5 Fig; S6 Table). The insensitivity of prr7-11 to NAM was confirmed by measuring circadian rhythms of leaf movement (S6 Fig). prr7-11 was not affected by NAM at any tested concentration, contrasting with a dose-dependent effect of NAM on circadian period in other prr mutants and associated backgrounds (R² > 0.9; Fig 5C).

In contrast, toc1-2 and TOC1-ox had significantly greater responses to NAM than wild type (Fig 5A; S6 Table). These results support our predictions that NAM induces dynamical changes specifically to PRR7 and TOC1. No dramatic changes of period, however, were observed for cry2-1 and cca1-11, suggesting that these might not contribute directly to the response to NAM.

Finally, derived from the nu-gap analysis, the possible change in the dynamical behavior of PRR9 in mediating the effect of NAM on the clock was evaluated with a prr7-3 and prr9-10 double mutant. prr7-3 and prr9-10 had an epistatic interaction, with the double mutant responding to NAM by a 5.3 ± 1.6 h increase of period, more than either the insensitive prr7-3 or the oversensitive prr9-10 alone (Fig 5A). The epistasis of prr9-10 to prr7-3 was confirmed at all concentrations of NAM tested (Fig 5C).

Nicotinamide-induced changes in period are associated with a blue light signaling pathway

The mutant analysis did not confirm the modeling dynamical perturbation of CRY2 in the response to NAM. However, CRY2 is one of a pair of cryptochrome blue light photoreceptors and so mutant analysis might not be the most appropriate tool to investigate the role of the blue light photoreceptor. To investigate further we also investigated the role of blue light in the response to NAM using monochromatic light conditions. High frequency measurements of the circadian promoter fusions PRR9::LUC, PRR7::LUC, TOC1::LUC, CCA1::LUC, LHY::LUC and
Fig 5. Reverse genetic analysis validates the prediction that TOC1 and PRR7 are associated with the effect of nicotinamide on circadian period. (A) The change in circadian period caused by 20 mM NAM (period difference) in 20

his circadian oscillator.
dyDE and the period control of the Arabidopsis circadian oscillator

DyDE and the period control of the Arabidopsis circadian oscillator

DyDE was further adapted to explore the rhythmic genome for additional targets for NAM and novel clock genes. For this purpose, models were computed between each pair of the 988 genes that were scored rhythmic in both untreated and NAM treated conditions, resulting in 2 million models corresponding to potential interactions.

We selected the models that exhibit the highest goodness of fit (over 80%) in both untreated and NAM-treated plants to minimize the identification of erroneous interactions and computed their nu-gap value to investigate dynamics affected by NAM. As a result, out of ten, two models only were retained with a nu-gap > 0.2. These models identified the regulation of AT5G35970 (P-loop containing nucleoside triphosphate hydrolases superfamily protein) by AT2G21860 (violaxanthin de-epoxidase-like protein) and the regulation of ATG21660 (GRP7/
CCR2) by AT1G78600 (LZF1/BBX22) as being altered by NAM. The regulation of AT5G35970 by AT2G21860 may be important as AT5G35970 is identified by DyDE as being a hub regulated by four circadian oscillator genes (S8 and S9 Tables). The second link is easier to explain because GRP7 along with GRP8 forms a slave oscillator driven by the circadian clock that regulates ABA responses [43]. GRP7 is an RNA binding protein regulated by ADP ribosylation [44]. As the enzymes that perform ADP ribosylation are inhibited by NAD, this could suggest

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a role for nicotinamide inhibiting ADP ribosylation of an oscillator or slave oscillator component.

Then, the fitness threshold was released to 60% to further investigate novel clock components. For this purpose, we searched for those genes for which models can be computed from/to clock components (S8 Table). Models with a nu-gap value above 0.2 were discarded as a consistency criterion. Finally, candidates were ranked according to their connectivity with the clock. As a result, 20 high potential genes were isolated (S9 Table). The whole genome analysis of clock input and output hubs and the nu-gap analysis suggest interesting roles for previously characterized genes, including AT3G47500 (CYCLING DOF FACTOR3) [45], AT4G38960 (BBX19) [46], AT1G78600 (BBX22) [47,48], AT3G22840 (CRY3) [49], AT1G28330 (DRM1), AT2G33830 (DRM2) [50,51] and uncharacterized genes including AT5G35970.

Discussion

The quantity and relevance of experimental measurements as well as a priori biological knowledge of the system are the two main factors that determine the choice of appropriate model complexity. Here, we considered the problem of inferring the entry point of a treatment in an organism from limited time-series data (in this case, the circadian clock in Arabidopsis). For this purpose, we used simple dynamical models to capture gene regulatory dynamics and compare those under different scenarios without making a priori assumptions on the structure of the network. Subsequently, we showed that simple dynamical models have the potential to identify crucial dynamical perturbations for complex systems such as the circadian clock. However, it should be stressed that, as for the sole purpose of identifying the topology of the underlying network, our method competes well with the current state-of-the-art of network inference strategies. To provide a more comprehensive picture, we further conducted a comparative analysis of the network reconstruction performances of our algorithm with four different methodologies [25,28,30,52] on a widely used circadian benchmark dataset generated from the Pokhilko 2010 model [25,53]. The simulations were carried to replicate the experimental conditions of our dataset (constant light, 48h of observational time, 4h sampling rate) [54,55]. S10 Fig and S10 Table displays the respective performance of each algorithm in terms of the resulting Area Under the ROC Curve and the Precision-Recall Curve. Precision-Recall curves allow for a more accurate picture of algorithms performances for sparse GRNs (such as the plant circadian clock) and are commonly used for such task. As a result, it can be seen that the performances of our methodology approach those of an adaptation of the best performer of the Dialogue for Reverse Engineering Assessments and Methods (DREAM) [26] to time-series data: the semi-parametric method dynGENIE3. The nonlinear parametric approach (GESBL) and the semi-mechanistic methodology (iCheMA) both seem to underperform. However, the non-parametric approach (GDPM) significantly outperformed all methods provided. This result is consistent with the currently available literature, which seems to favour the use of nonparametric nonlinear equations for the inference of network topology from short-time series data. However, as mentioned above, comparison of nonlinear dynamics is not trivial, and require further mathematical developments. Hence, as a future work, it would be interesting to address the development of such suitable mathematical tools.

We further devised a learning algorithm to capture the specific pattern of oscillating wave forms of genes affected by NAM. Since the period of oscillations of central clock genes increases from 24 (wildtype) to roughly 28 (NAM) hours, we focused on those genes. For a relatively small number of genes, DyDE efficiently narrowed down possible targets of NAM that could then be verified experimentally. Since it is likely that other genes may be targets of
NAM, we further applied DyDE to all 988 circadian genes that were scored rhythmic in both untreated and NAM treated conditions.

DyDE identified important changes in the regulatory dynamics of PRR7, TOC1, CCA1 and the blue light photoreceptor, CRY2, resulting from the treatment of plants to NAM as well as suggesting a mediating role of PRR9. Mutants analysis confirmed DyDE predictions of altered activity of PRR7, TOC1 and PRR9 and blue/red light experiments demonstrated that the effect of NAM is blue light dependent. The latter also demonstrated that blue light regulates circadian oscillations of $[\text{Ca}^{2+}]_{\text{cyt}}$ through a NAM-sensitive pathway.

The involvement of PRR7 with the dynamic adjustment of circadian period in response to nicotinamide, revealed by the insensitivity of prr7-11 and prr7-3 to NAM and confirmed by leaf movements analysis, is interesting because PRR7 is also required for the response of the circadian oscillator to sugars [10,56]. PRR7, however, is not a direct target for NAM in the circadian oscillator because PRR7 is not required for the response to NAM, as demonstrated by the hyper-sensitivity to NAM of the prr7-3 prr9-10 double mutant. Together, the insensitivity of prr7-3 and prr7-11 to NAM and hypersensitivity in the prr7-3 prr9-10 double mutant indicates that PRR7 and PRR9 regulate a component or pathway influenced by NAM and that PRR7 might act upstream of PRR9 in this regulation. The levels of expression of PRR7 and 9 appear to regulate the pace of the circadian oscillator through feedback with CCA1/LHY and by acting as toggle switching the oscillator from a morning state of CCA1/LHY are high to an evening state of TOC1 is high [57,58].

Additionally, the blue-light dependency of both circadian oscillations of $[\text{Ca}^{2+}]_{\text{cyt}}$ and NAM regulation of circadian period might suggest that $\text{Ca}^{2+}$ is associated with the response of the oscillator to NAM. Furthermore, we recently reported that CALMODULIN-LIKE 24 (CML24), is a $\text{Ca}^{2+}$-dependent regulator of circadian period and that its affects are NAM sensitive [59]. A caveat to this argument is that our methodology identified CRY2 regulation of the transcriptional network being altered by NAM but the circadian oscillations of $[\text{Ca}^{2+}]_{\text{cyt}}$ were dependent on CRY1. NAM can also affect the oscillator through $\text{Ca}^{2+}$-independent mechanisms [14]. We propose that a module of circadian oscillator components PRR7 and 9, TOC1 and a $\text{Ca}^{2+}$ signaling network contribute to the blue light-dependent response of the circadian oscillator to NAM that regulates circadian period (Fig 7).

Then, extension of DyDE to the whole circadian genome has also identified components outside the core oscillator that might also be involved in response to NAM, including the regulation of GRP7/CCR2 by LZF1/BBX22 and theses will be candidates for future investigation. Remarkably, five genes out of 22 that were isolated in our genome analysis are known to interact with circadian regulators (BX19, CYCLING DOF FACTOR3 [45,46], have been previously implicated in circadian regulation (BBX22) [47,48], in blue light signaling (CRY3) [49] or are being downregulated by ABA/cADPR (GRP7, BBX19, BBX22) [43,44,60]. This result is encouraging and opens the door to the identification of novel drivers of circadian rhythms in Arabidopsis.

Overall, we suggest that the description of gene regulatory dependencies and the quantification of changes in dynamics computed by DyDE provide reliable hypotheses for the investigation of drug targets in complex gene regulatory networks, which has a broad range of applications in systems biology.

**Experimental procedures**

**Plant material and growth conditions.** Arabidopsis thaliana C24 ecotype untreated plant carrying pCHLOROPHYLL A/B BINDING-PROTEIN2:LUCIFERASE (CAB2:LUC) and p35SCAMV:AEOQUORIN (35S:AEQ) and photoreceptors mutants carrying 35S:AEQ were
previously reported [61]. All other plant lines were previously reported [10]. Prr7-3 and prr7-
11 are the same T-DNA insertion loss-of-function mutation being independently character-
ised as prr7-3 [62] and prr7-11 [63]. These were maintained as separate populations. Growth
conditions were as previously described [10].

**Microarray datasets.** C24 seeds were grown for 10 days. On the 11th day the seedlings
were exposed to constant light (70 μmol m⁻² s⁻¹). 50% of C24 plants were treated with 50 mM
NAM (Sigma, U.K.) every two hours as previously described [41]. Aerial tissue from 100 seed-
lings was sampled every 4 hours from ZT 49 to 93. RNA extraction was performed as described
[12]. Three independent experiments were carried out and 100 ug of RNA from two of these
was sent to NASC (Nottingham, UK) for microarray analysis using the ATH1 array. Note
that PRR9 and LUX probes on the ATH1 microarray also measure the expression levels of
AT2G46670 and AT5G59570, respectively. Comparing other datasets [40] shows that the
probe with PRR9 agrees with PRR9 on other datasets while the one with LUX does not. Hence,
we used this probe to represent PRR9, but not LUX.

All modeling and analysis was carried out in MATLAB unless stated otherwise. Microarray
data was normalised using gcRMA. Go enrichment was determined using Gorilla [64]. Pro-
moter motif enrichment was calculated using AtCOECIS [65].
Validation of microarray data sets. The microarrays were validated by RT–qPCR using aliquots of the RNA extracted for microarray hybridization as described previously [10]. CCA1 and TOC1 were analysed and quantified as detailed previously [10].

Luminescence time course experiments. Photon counting imaging of aequorin or luciferase activity in constant light of 60–70 μmol m⁻² s⁻¹ monochromatic blue, monochromatic red or equal blue-red LED light was performed as described [12]. Circadian period estimates were calculated using the MS Excel BRASS plugin (http://amillar.org/downloads).

Supporting information

S1 Fig. Detrended time-series of circadian transcript in both untreated and NAM condition. Data were gathered for 44 hours every 4 hours, 2 replicates, starting from 49 hours after the switch to constant light (i.e., third day of constant light). Data showed are detrended, so that the rhythmic pattern is clear. LUX does not appear on this list, as the probe also measured the expression of AT5G59570. CRY1, PHYB, ELF3, ZTL and CHE were not considered for the network inference step.

S2 Fig. Assessment of circadian regulated transcripts from both the learning methodology and standard tools. (A) Results correspond to untreated plants. The trained algorithm, COSOPT [33] and JTK [32] respectively identified 3859, 1856 and 3698 circadian regulated transcripts. JTK and the trained algorithm identified most of the genes labelled as periodic by COSOPT (resp. 87% and 81% of them). The rhythmicity of 75% of the genes labelled rhythmic by JTK was confirmed by the learning strategy. (B) Results corresponding to NAM-treated plants. The rhythmicity of 60% of the genes labelled rhythmic by JTK was confirmed by the learning strategy while 1636 novel genes were identified as rhythmic with a typical non-sinusoidal profile.

S3 Fig. Coverage and false positive curves of the known regulatory links involved in the circadian oscillator of Arabidopsis Thaliana, as inferred by DyDE. In DyDE, linear Ordinary Differential equations (ODEs) of order one are computed between each pair of genes to describe the dynamics of the whole system. To be further considered as a good approximation of the dynamics involved, each dynamical model needs to pass a validation criterion based on its agreement to the data (i.e. a user-defined threshold on the goodness of fit). Decreasing the fitness threshold leads to a better coverage (upper panel) of the system dynamics but increase the amount of false positives (lower panel). The coverage describes the amount of links inferred over the amount of total true links in the system (as defined by Fogelmark et al, 2014). The number of false positive corresponds to links that are not represented in Fogelmark et al. The maximum amount of possible false positives is 32, while the total amount of links in the true system is of 40. The threshold of 46% (represented by a red cross) is chosen for this analysis with a coverage of 70% (which corresponds to 28 true positives and 21 false positives).

S4 Fig. Networks inferred by DyDE in both untreated and NAM-treated plants, with the regulation loss networks. (A) Network inferred in absence of NAM. (B) Network inferred in presence of NAM. (C) Regulation loss network. (D) Common regulation network. On these graphs, genes are organized by their peaking time during diel cycles. Genes that are represented by a yellow filled circle peak in the morning, dark brown filled circles stand for an evening peaking time and grey for a night peaking time. White filled circles correspond to photoreceptors. Networks are made of the conjunction of identified LTI models for which
respective fitness values are above 46%. Blue arrows correspond to activation whereas red arrows correspond to an identified inhibition (S3 Table). On (D), the highlighted links represent the models with the five highest nu-gap values.

EPS

S5 Fig. The effect of nicotinamide on circadian rhythms in circadian, light and ABA signaling mutants. Rhythms of luciferase activity were measured in the presence (grey) or absence (white) of 20 mM NAM. Col-0 CCA1:LUC, Col-0 TOC1:LUC, WS CAB:LUC⁺ and C24 CAB: LUC backgrounds are shown with respective mutants. Traces represent mean values of n > 16 biological replicates from > 2 technical replicates in the presence (grey) or absence (clear) of 20 mM NAM.

EPS

S6 Fig. Circadian rhythms of leaf movement are insensitive to nicotinamide in prr7-11. Rhythms of leaf movement were measured in constant white light for 120 h in the presence (yellow) or absence (white) of 20 mM NAM. n = 22 Col-0, and n = 30 prr7-11.

EPS

S7 Fig. Circadian rhythms of CCA1:LUC, LHY:LUC, PRR7:LUC, PRR9:LUC, TOC1:LUC and GL1:LUC activity in constant red and blue mixed light, constant red or constant blue light (70 umol m⁻² s⁻¹). Grey indicates the presence and white indicates the absence of 20 mM NAM. Light conditions indicated by the colored boxes on the X axes. White is red/blue mix, monochromatic light is indicated by the appropriate color with subjective night shaded darker than subjective day.

EPS

S8 Fig. Phytochrome and cryptochrome modulate [Ca²⁺]_cyt under monochromatic light. 35S:AEQ luminescence measured over 96 hours in constant 65 umol m⁻² s⁻¹ monochromatic red or blue light in light signaling mutants. Bars on the X axis indicate red and blue mixed light (white), dark (black), monochromatic red (red), or monochromatic blue (blue). Darker boxes indicate subjective night in constant light. N = 8. Error bars SEM.

EPS

S9 Fig. Blue light-sensitive circadian [Ca²⁺]_cyt oscillations are inhibited by nicotinamide. 35S:AEQ luminescence measured over 96 hours in constant 65 umol m⁻² s⁻¹ monochromatic red or blue light in light signaling mutants in the presence (grey circles) and absence (open circles) of 20 mM NAM. Bars on the X axis indicate light treatment of red and blue mixed light (white), dark (black), monochromatic red (red) or monochromatic blue (blue). Darker boxes indicate subjective night in constant light. n = 8. Error bars SEM.

EPS

S10 Fig. Comparative analysis of the network reconstruction accuracy of our algorithm with state-of-the-art network inference methodologies. The Area Under the ROC Curve and the Precision-Recall Curve are shown for each algorithm and simulation run. For this comparison, only 10 stochastic simulations were carried to limit the overall computation time of methodologies that involve a Markov Chain Monte Carlo (MCMC) sampling scheme. Each box on these graphs represents the 50th – 95th percentile around the median. Dynamical GENIE3 (dynGENIE3, [28]) is an adaptation of the GENIE3 [51] method for time series data, which was the best performer of the DREAM4 Multifactorial Network challenge and the DREAM5 Network Inference Challenge [26,22]. Like our approach, the temporal evolution of genes in dynGENIE3 is based on an ODE model. However, in this specific case, the transcription function in each ODE is represented by ensembles of regression trees and is therefore not
fixed a priori. This semi-parametric approach provides a greater flexibility to the inference framework but complicates the comparison of dynamical properties between experimental conditions. In contrast, Continuous-time Gaussian process dynamical model (GPDM) is a non-parametric, Gaussian process-based GRN inference algorithm for which the performances were compared to the best performers of the DREAM challenge and consistently shown superior in dealing with short time series data generated from the 10-genes challenge [30]. A parametric approach, so-called Sparse Bayesian Learning (GSBL) has also been investigated to identify a sparse representation of GRN topology with nonlinear multivariate ARX models and showed promising performances on a repressilator model of gene regulation [52]. Finally, a semi-mechanistic model based on gradient matching and nonlinear Hill-type transcription function such as the iCheMA [25] algorithm have been developed and was revealed as the best performer of a set of established state-of-the-art network reconstruction methods applied to the inference of circadian-type regulatory networks. Our algorithm reached an AUROC of 0.6426 +- 0.0448 and an AUPREC of 0.5124 +- 0.049. dynGENIE3 obtained an AUROC of 0.6709 +- 0.0427 and an AUPREC of 0.512 + 0.049. GPDM obtained an AUROC of 0.7119 +- 0.0538 and an AUPREC of 0.6759 + 0.0745. GSBL obtained an AUROC of 0.5801 + 0.0542 and an AUPREC of 0.4999 + 0.707 while iCheMA obtained an AUROC of 0.5523 + 0.0874 and an AUPREC of 0.2848 + 0.0397. Details of the results for each run are presented on the S10 Table.

S1 Table. Circadian transcriptomes of untreated and nicotinamide-treated plants. Circadian time-series for the transcripts on the ATH1 microarray for the untreated and nicotinamide-treated (NAM) datasets are presented on three separate sheets. Gene lists are provided describing the genes that were identified by PEAL to be circadian-regulated, along with the estimated period, amplitude and phase, and confidence of estimation for C24, NAM and toc1-1. The final two sheets describe the genes common between the C24 and NAM, and the C24 and toc1-1 circadian-regulated subsets.

S2 Table. GO enrichment terms of transcripts rhythmic. The tables list the most enriched terms in the rhythmic transcripts, split into the three ontologies: Biological process, Molecular function, and Cellular component. The analysis is split in eight categories, each corresponding to a column in the table: GO term (unique identifier in the GO database), GO description, p-value (likelihood of enrichment), False Discovery Rate q-value, a measure of Enrichment, a link to the full list of enriched genes, and whether the GO term is also enriched in the NAM datasets.

S3 Table. Details of the dynamical interactions inferred by DyDE between the genes of the Arabidopsis thaliana circadian oscillator. The tables describe the connections modelled for the circadian oscillator in the untreated plants, NAM treated, regulation loss and common networks. Network connections are described gene by gene and a comparison with the known regulatory interactions is provided. The fitness value is displayed, as well as the sign of the regulation (activation or inhibition).

S4 Table. Sorted v-gap values corresponding to common links between untreated and NAM. Nu-gaps values computed for each link inferred in both untreated and NAM-treated networks. This table ranks the nu-gap values from the largest to the smallest.
S5 Table. Connectivity loss corresponding to each gene, for untreated and NAM-treated networks. Values are displayed for a fitness threshold of 46%.

S6 Table. The effect of nicotinamide on circadian period in circadian oscillator mutants. The period of circadian rhythms of luciferase activity were measured in the presence absence of 20 mM NAM in Col-0 CCA1:LUC, Col-0 TOC1:LUC, WS CAB:LUC* and C24. CAB:LUC backgrounds are reported with the respective mutants. n > 16 biological replicates from > 2 technical replicates.

S7 Table. Cost functions and parameters optimization of the P2012 model. These tables sort the cost function scores obtained by sequentially optimizing a single and two parameters simultaneously to the best fit of TOC1 and CCA1 time-series profile in the presence of NAM. Best performances for single parameter tuning are achieved for m14 and m15 respectively while the best performances for two simultaneous parameters are obtained through a mixture of LHY and PRR7 parameters.

S8 Table. Extension of the DyDE methodology to the whole genome: List of potential novel candidates involved in clock regulation. The list of models that have been validated (> 60% fitness, nu-gap < 0.2) between the genome and the known clock genes. The lists are split in four sheets, displaying separately the analysis of the genome to the main clock genes and vice versa. The genes are sorted in alphabetical order. Their function and the sign of the identified regulation are provided, as well as the count of appearance of each gene in their respective list.

S9 Table. Summary of the novel potential clock genes identified. The list of genes identified as input/output hubs to the clock as well as their functions is represented together with their potential regulation by cADPR.

S10 Table. Details of the performances of state-of-the-art network inference methodologies against our approach. This table summarize the Area Under the ROC Curve and the Precision-Recall curve obtained by each algorithm, for each run. Highlighted is the best performer for each run.

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