**DREISS: dynamics of gene expression driven by external and internal regulatory networks based on state space model**

Daifeng Wang1,2, Fei He4, Sergei Maslov4, Mark Gerstein1,2,3\*

1Program in Computational Biology and Bioinformatics; 2Department of Molecular Biophysics and Biochemistry; 3Department of Computer Science, Yale University, New Haven, CT, USA. 4Biological, Environmental and Climate Sciences Department, Brookhaven National Laboratory, Upton, NY, USA. \*Correspondence to: [pi@gersteinlab.org](mailto:pi@gersteinlab.org)

[[1]](#footnote-2)\*abstract

Gene expression is controlled by combinatorial effects of gene regulatory factors from different biological subsystems driving specific regulatory functions such as general transcription factors, cellular growth factors and microRNAs. A subsystem’s gene expression may be controlled by its internal regulatory factors, exclusively, or by other external subsystems, or by both. It is thus useful to distinguish the degree to which a subsystem is regulated internally or externally; e.g., how external regulatory factors affect the expression of conserved genes during evolution.

We developed a computational method, DREISS for dynamics of gene expression driven by external and internal regulatory modules based on state space model to help dissect the effects of different regulatory subsystems on gene expression (https://github.com/gersteinlab/Dreiss). Given a subsystem, the “state” and “control” in the model refer to its own (internal) and another subsystem’s (external) gene expression levels. The state at a time is determined by the state and control at previous time. DREISS uses the dimensionality reduction to combat the limited time samples, and identifies the canonical temporal expression trajectories (e.g., degradation, growth, oscillation) representing the regulatory effects from various subsystems.

To illustrate DRIESS, we applied it to the time-series gene expression datasets of *C. elegans* and *D. melanogaster* during their embryonic development, to demonstrate its capabilities for studying the regulatory effects of evolutionary conserved vs. divergent transcription factors across distant species. We analyzed the expression dynamics of the conserved, orthologous genes (orthologs), seeing the degree to which these can be accounted for by orthologous (internal) versus species-specific (external) transcription factors (TFs). We found that between the two species, the orthologs canonical expression trajectories driven by orthologous TFs are more similar to each other than those driven by species-specific ones. This is particularly true for genes with evolutionarily ancient functions (e.g. the ribosome), in contrast to those with more recently evolved functions (e.g., cell-cell communication). This implies that despite striking morphological differences, some fundamental embryonic-developmental processes are still tightly under the control of an ancient regulatory system for the similar canonical trajectories of worm-fly orthologs driven by the orthologous TFs.

# introduction

Gene regulatory networks (GRNs) systematically control the gene expression dynamics. These networks are highly modular, and consist of various sub-networks. Each sub-network contains a number of regulatory factors representing a subsystem that drives specific gene regulatory functions [[1](#_ENREF_1),[2](#_ENREF_2)]. The subsystems interact with one another, and work together to carry out the entire gene regulatory function. For example, the gene expression in embryogenesis is controlled by the combinatorial effects of various regulatory subsystems composed of complex evolutionary regulatory networks [[3](#_ENREF_3)]. These regulatory subsystems drive very diverse developmental programs, from the highly conserved (e.g. DNA replication) to the species-specific (e.g. body segmentation). As such the orthologous genes that are evolutionary conserved genes across species can therefore be regulated by both orthologous and species-specific transcription factors (TFs) [[4](#_ENREF_4)]. The orthologous TFs form an “internal” regulatory network, while the species-specific TFs form an “external” one. Unfortunately, existing experimental gene expression data cannot decouple the expression components that are driven by the different subsystems. Thus, computational methods are required to assess the contribution from each factor or subsystem from the gene expression data. In this study, we propose a novel computational method, DREISS - dynamics of gene expression driven by external and internal regulatory networks based on state space model. Using DREISS, we are able to identify temporal gene expression dynamic patterns for evolutionarily conserved genes during embryonic development, as driven by conserved and species-specific regulatory subsystems. These results advance our current understanding of GRNs during evolution, as well as the differentiation during development.

Developmental GRNs control gene expression during the developmental processes. These particular GRNs have evolved, making it difficult to understand their regulatory mechanisms at the system level. Hence, one typically compares developmental gene expression across species to infer biological activities of developmental GRNs. For example, embryogenesis provides a platform to study the evolution of gene expression between different species. Recent work has showed that significant biological insight can be gained by cross-species comparisons of the expression profiles during embryogenesis for worms [[5](#_ENREF_5)], flies [[6](#_ENREF_6)], frogs [[7](#_ENREF_7)] and several other vertebrates [[8](#_ENREF_8)]. It was found that the orthologous genes have minimal temporal expression divergence during the phylotypic stage, a middle phase during the embryonic development across species within the same phylum. These patterns are often characterized as “hourglass” [[9](#_ENREF_9)]. In addition, the conserved hourglass patterns were observed even within a single species while comparing the developmental gene expression data across distant species, such as worm and fly [[10](#_ENREF_10)]; i.e., the expression divergence among evolutionarily conserved genes become minimal during the phylotypic stage in both worm and fly. However, much less is known about how the orthologous genes in each species eventually contribute to their species-specific phenotypes due to the lack of appropriate computational approaches. Thus, we aim to use DREISS to discover the components of the orthologous gene expression during embryonic development driven by species-specific transcription factors.

The state-space model has been widely used in engineering [[11](#_ENREF_11)], and also in biology for the analysis of gene expression dynamics [[12-14](#_ENREF_12)]. It models the dynamical system output as a function of both the current internal system state and the external input signal. A commonly used example in engineering is the vehicle cruise control system where the system output and state is the vehicle’s speed. Based on the road conditions, the cruise control requires various fuel amounts in order to keep the desired speed level. In biology, we can look at the transcription factors and microRNAs as internal and respectively external regulatory factors of the protein-coding genes expression (See more internal-external examples in Supplemental Table 1). Similarly, the state-space model can be applied for studying the expression of orthologous genes at different developmental stages using information regarding their expression (internal) and species-specific regulatory factors (external) at the current known developmental stage. Unlike earlier studies that calculate the expression correlation between individual genes, the state-space model predicts the temporal causal relationships at the system level; i.e., the state at a time is determined by the state and external input at the previous time. The earlier work applied the state-space model to study the gene expression dynamics focusing on small-scale systems, and did not explore the analytic dynamic characteristics of the inferred state-space models. The complex and large-scale biological datasets, especially temporal gene expression data, are very noisy, and high dimensional (i.e., the number of genes is much greater than the number of time samples), thereby preventing an accurate estimation of the state-space model’s parameters. The dimensionality reduction techniques have thus been used to project high-dimensional genes to low-dimensional meta-genes (i.e., the selected features representing de-noised and systematic expression patterns [[1](#_ENREF_1),[15](#_ENREF_15),[16](#_ENREF_16)]) as well as the principal dynamic patterns for those meta-genes [[17](#_ENREF_17),[18](#_ENREF_18)]. Using DREISS, we are able to apply the dimensionality reduction to the gene expression data, and develop an effective state-space model for their meta-genes, and then identify a group of canonical temporal expression trajectories representing the dynamic patterns driven by the effective conserved and species-specific meta-gene regulatory networks according to the model’s analytic characteristics. These dynamic patterns reveal temporal gene expression components that are controlled by conserved or species-specific GRNs.

DREISS is a general-purpose tool and can be used to study the gene regulatory effects from any different subsystems for a given group of genes. As an illustration, we applied DREISS to the gene expression data during embryonic development for two model organisms, worm (*Caenorhabditis elegans*) and fly (*Drosophila melanogaster*). In both species, we were able to identify the expression patterns of worm-fly orthologs driven by the conserved regulatory network consisting of the worm-fly TFs (i.e., the conserved regulatory subsystems between two species), as well as the worm/fly-specific regulatory network consisting of non-orthologous TFs (i.e., the species-specific regulatory subsystem). Our results reveal that, in addition to executing conserved developmental functions between worm and fly, their orthologous genes are also regulated by species-specific TFs to involve in species-specific developmental processes. In summary, DRIESS provides a framework to analyze both distantly and closely related species allowing for a better understanding of the gene regulatory mechanisms during development.

# methods

DREISS consists of five major steps as detailed in Figure 1:

**Step A**: DREISS models temporal gene expression dynamics using state-space models in control theory. In this step, we need to define the internal and external groups of genes and input their time-series gene expression data that we are interested to study. We assume that the time-series gene expression data fits a state-space module. In the state-space model, the “state” refers to the expressions for a large group of genes of interest, such as the worm-fly orthologous genes investigated here. The “control” refers to any other group of genes that contribute to gene expressions of the “state”, such as the species-specific TFs contributed to control orthologous gene expression.

**Step B**: Due to the limited number of temporal samples in gene expression experiments, we do not have enough data to accurately estimate the parameters of the state-space models that capture interactions among hundreds of genes. Therefore, DREISS projects high-dimensional gene expression space to lower-dimensional meta-gene expression spaces using dimensionality reduction techniques.

**Step C**: DREISS then derives the effective state-space models for meta-genes so that model parameters can be estimated.

**Step D**: DREISS then identifies the meta-gene expression dynamic patterns; i.e., canonical temporal expression trajectories driven by “state” (internal) and by “control” (external) based on the analytic solutions of the estimated models.

**Step E**: Finally, DREISS calculates the coefficients of genes for the dynamic patterns of linear transformations between genes and meta-genes. DREISS also allows us to compare the dynamic expression patterns of multiple datasets with samples taken at different times. We describe each DREISS step in detail as follows.

## State-space models for temporal gene expression dynamics

A gene regulatory network is made up of various subsystems [[1](#_ENREF_1),[2](#_ENREF_2)]. These subsystems work together to execute the regulatory functions. Given a group of *N*1 genes in a subsystem, their gene expression levels (*X*) are not only controlled by internal interactions among *X*, but also affected by the regulatory factors from other subsystems outside *X* (external regulations denoted here by the *U* group). For example, we consider the worm-fly orthologous genes as the *X* group. The worm-fly orthologous TFs from the *X* group are the *internal* regulatory factors, and non-orthologous TFs such as worm- or fly- specific TFs are the *external* regulatory factors to the *X* group, namely Group *U*. Both the internal and external regulatory factors control gene expressions in dynamic ways (i.e., their regulatory signals at the current time will affect gene expressions at subsequent times). Thus, the regulatory mechanisms for the gene expressions form a control system. In this study, we used a state-space model (defined by linear first-order difference equations, Figure 2A) to formulate temporal gene expression dynamics for the gene group *X* (comprising *N*1 genes) with external regulation parameters from the gene group *U* (comprising *N*2 genes) at time points *1*, *2*, … , *T* as follows:

(1)



, where the vector , the “state”, includes *N*1 gene expression levels at time *t* in group *X*, and the vector , the “input or control”, includes *N*2 gene expression levels at time *t* in group *U*. The system matrix captures internal causal interactions among genes in *X* (i.e., the *i*th, *j*th element of *A*, *Aij* describes the contribution from the *j*th gene expression at time *t* to the *i*th gene expression at the next time *t*+1), which instantiates a gene regulatory network. The control matrix captures external causal regulations from the genes in *U* to genes in *X* (i.e., the *i*th, *j*th element of *B*, *Bij* describes the contribution from the *j*th gene expression in *U* at time *t* to the *i*th gene expression in *X* at the next time *t*+1). represents the real number domain. According to the state space model (1), the gene expression dynamics in *X* is determined by the system matrix *A* and the control matrix *B*.



## Dimensionality reduction from genes to meta-genes

The temporal gene expression experiments normally have limited time samples (for example, there may only be a dozen time points), which are far less than the time samples needed to estimate the large matrices *A* and *B* when *X* and *U* are composed of hundreds or thousands of genes. Thus, we project high dimensional temporal gene expressions to much lower dimensional meta-gene expression levels using a dimensionality reduction technique (Figure 2B). Those meta-gene expression levels should capture original gene expression patterns, such as the ones having the greatest degree of co-variation. We calculate the meta-gene expression levels as follows:

(2)

, where , the “meta-gene state” at time *t*, includes *M*1 (<< *N*1 and <*T*) meta-gene expression levels; i.e., the first *M*1 elements of the *t*th row of the matrix whose columns are right-singular vectors of the matrix in group *X* by the singular value decomposition (SVD) [[19](#_ENREF_19)]; the vector , the “meta-gene input or control” at time *t*, includes *M*2 (<< *N*2 and <*T*)meta-gene expression levels; i.e., the first *M*2 elements of the *t*th row of the matrix whose columns are right-singular vectors from SVD of the matrix in group *U*; is the linear projection matrix of SVD from *M*1 meta-gene expression space to *N*1 gene expression space in *X,*  is the linear projection matrix of SVD from *M*2 meta-gene expression space to *N*2 gene expression space in *U*, and (.)\* is a pseudo-inverse operation; i.e., *W\*W*=*I*, where *I* is the identity matrix.



## Estimation of effective state-space model for meta-gene expression dynamics

Next, we obtain the effective state-space model for meta-genes using linear projections *WX* and *WU* between genes and meta-genes as follows (Figure 2C). By replacing (1) using (2), we obtain that

(3)



, and by multiplying the pseudo-inverse of *WX*, where *I* is an identity matrix, at both sides of (3),



(4)



, where the effective meta-gene system matrix captures internal causal interactions among meta-genes in *X* (i.e., the *i*th, *j*th element of () describes the contribution from the *j*th meta-gene expression at time *t* to *i*th meta-gene expression at time *t*+1), and the effective control matrix captures external causal regulations from meta-genes in *U* to meta-genes in *X* (i.e., the *i*th, *j*th element of , describes the contribution from the *j*th meta-gene expression in *U* at time *t* to *i*th meta-gene expression in *X* at time *t*+1). Equation (4) describes the effective state space model for the meta-genes in X, whose expression dynamics is determined byand. Because the meta-gene dimension, *M*1 (*M*2) is less than *T*, and much less than *N*1 (*N*2), we can estimateand as follows.



We rewrite Equation (4) as a matrix product on the right side:

. (5)



By applying Equation (5) to time points, 2,3, … , *T*, we then obtain that

(6)



, where .



The effective internal system matrix and external control matrixcan be estimated by:



(7)



, where is the pseudo-inverse of ; i.e.



## Identification of internally and externally driven principal dynamic expression patterns of meta-genes (canonical temporal expression trajectories)

The analytic solution to a general first-order linear matrix difference equation [[20](#_ENREF_20)], *Yt*+1=*CYt* is

*Yt*=*CtY0=(PDP-1) t Y0=PDtP*-1*Y0=PDtS*, where the columns of the matrix *P* are eigenvectors of *C*, the diagonal elements of the diagonal matrix *D* are eigenvalues of *C* such that *CP*=*PD*, and the vector *S*= *P*-1*Y0*. Then, if we rewrite *Yt* by a linear combination of the time exponential of eigenvalues of *C*, we have that , where *mc* is the total number of eigenvalues of *C*, *αi* is the *i*th eigenvalue of *C*, *si* is the *i*th element of *S*, *Pi* is the *i*th eigenvector of *C* (i.e., the *i*th column of *P*), and *Ki*=*siPi* is the coefficient vector of *Yt* over the *t*th time exponential of *αi*.

From Equation (4), the internally driven components of meta-gene states at two adjacent time points have . According to the above analytic solution, the components of meta-gene expressions in *X* driven by effective internal regulations are linear combinations of *M*1 dynamic patterns determined by the eigenvalues of the effective system matrix as follows:



; i.e., the internally driven component of *i*th meta-gene’s expression across all time points, (8)

, where and are the *p*th eigenvalue of and its coefficient vector from the analytic solution, which determines the *p*th dynamic pattern driven by effective internal regulations, defined as the *p*th internal principal dynamic pattern (iPDP) = , in which represents the *t*th power of , and *V*(*i*) represents *i*th element of the vector *V*. represents the complex number domain, and “(*i*)” represents the *i*th element of vector. If an eigenvalue *λ* is complex when is asymmetric, then its conjugate is also an eigenvalue, so we sum its iPDP and its conjugate eigenvalue, ’s iPDP, as a unified iPDP with real elements equal to .

Similarly, the components of meta-gene expressions in *X* driven by effective external regulations from *U*, i.e., (externally driven components of meta-gene states at two adjacent time points) are linear combinations of *M*2 dynamic patterns determined by the eigenvalues of the effective system matrix as follows:



; i.e., the externally driven component of *i*th meta-gene’s expression across all time points, (9)

, where and are the *q*th eigenvalue of and its coefficient vector, which determines *q*th dynamic pattern driven by effective external regulations, defined as *q*th external principal dynamic pattern (ePDP) = , in which represents the *t*th power of , and *V*(*i*) represents *i*th element of the vector *V*. If an eigenvalue is complex, then its conjugate is also an eigenvalue, so we sum its ePDP and its conjugate eigenvalue, ’s ePDP, as a unified ePDP with real elements equal to .

Both the internal and external principal dynamic patterns (PDPs) represent the canonical temporal expression trajectories, which can be either increasing, or damped oscillation and so on depending on PDP’s eigenvalues (Table 1).

**Table 1**. Classification of canonical temporal expression trajectories for PDP eigenvalue types

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| PDP  eigenvalue | Real | | | | | | Complex (radius) | | |
| >1 | =1 | <1 & >0 | <0 & > -1 | = -1 | <-1 | >1 | =1 | <1 |
| Canonical temporal expression trajectory (initial) | Increasing (I) | Flat (F) | Decreasing (D) | Vibrating early (VE) | Vibrating (V) | Vibrating late (VL) | Underdamped oscillation (UO) | Oscillation (O) | Damped oscillation (DO) |

## Identification of gene coefficients of principal expression dynamic patterns

Because genes and meta-genes have linear relationships in terms of their expression levels as described in Equation (2), the components of gene expression levels in *X* driven by internal regulations, can be also expressed as linear combinations of *M*1 iPDPs:

; i.e.,

the internally driven component of *i*th gene’s expression across all time points, (10)

, where represents the gene coefficient vector for *p*th iPDP. Similarly, the gene expression components driven by external regulations from *U*, can be also expressed as linear combinations of *M*2 ePDPs:

; i.e.,

the externally driven component of *i*th gene’s expression across all time points, (11)

, where represents the gene coefficient vector for *q*th ePDP.

# results

Gene expression data during embryogenesis provide important information about the dynamics of genomic functions throughout the developmental process, from the conserved functions such as DNA replication to the species-specific functions such as body segmentations, but hardly reveal any data regarding the evolutionary gene regulatory subsystems that drive those developmental functions [[3](#_ENREF_3)]. Thus, in order to understand the relationships between those subsystems and their driving genomic functions, we apply DREISS to worm and fly gene expression datasets during embryogenesis in modENCODE and we are able to identify various developmental genomic functions of worm-fly orthologous gene pairs driven by two different evolutionary regulatory subsystems, conserved (worm-fly orthologous TFs) and non-conserved (worm/fly specific TFs). As model organisms for developmental biology, both worm and fly have been used previously to study embryogenesis.

## Applications to worm and fly embryonic developmental data in modENCODE: orthologous genes, transcription factors and gene expression datasets

DREISS enables us to compare expression dynamic patterns between two or more temporal gene expression datasets even though they have different numbers of samples, as well as differences in the times at which those samples were collected. For example, we can apply DREISS to two different datasets of the same group of genes, and identify both the common (similar) and the specific (different) dynamic patterns driven by internal regulations captured by the eigenvalues of the effective system matrices between the two datasets.

In this paper, we apply DREISS to 3,153 one-to-one orthologous genes between worm (*Caenorhabditis elegans*) and fly (*Drosophila melanogaster*) as Group *X* to study their expression dynamics during embryonic development [[10](#_ENREF_10)]. We refer to species-specific TFs as external regulations; i.e., Group *U*. We found that worm-fly orthologs have similar internal dynamic patterns, which may be mainly driven by conserved TFs, but have very different external dynamic patterns driven by species-specific TFs between worm and fly embryonic developmental stages. The data is summarized as follows.

We define Group *X* as 3,153 one-to-one orthologous genes between worm and fly during embryonic development, and Group *U* as all the species-specific TFs (509 worm-specific TFs, 442 fly-specific TFs) [[21](#_ENREF_21),[22](#_ENREF_22)]. We used their temporal gene expression levels (as measured by the RPKM values in RNA-seq) during embryonic development from the modENCODE project [[10](#_ENREF_10)]. The worm embryonic development dataset includes *T*=25 time stages at 0, 0.5, 1, 1.5, … , 12 hours, and the fly dataset includes *T*=12 time stages at 0, 2, 4, … , 22 hours, but *t*=1,2,..,25 for worm and *t*=1,2,…,12 for fly are used in this paper, representing the relative time points for the entire embryonic development processes. Because *M*1+ *M*2<*T* in Equation (7), we choose *M*1= *M*2=5 meta-genes for fly (*T*=12), and find that five meta-genes of Group *X* and five meta-genes of Group *U* capture ~98% of the co-variation of orthologous gene expressions and fly-specific TF gene expressions, respectively. In order to compare worm and fly, we also choose *M*1=*M*2=5 meta-genes for worm, which capture ~98% of the co-variation of orthologous gene expressions and worm-specific TF gene expressions.

## Meta-genes of worm-fly orthologous genes have similar internal, yet different external principal dynamic patterns during embryonic development

We find that the meta-gene canonical temporal expression trajectories driven by conserved regulatory networks (i.e., internal principal dynamic patterns, iPDPs) include four major patterns in both the worm and fly embryonic developmental process by order of eigenvalues: 1) a late highly varied pattern; 2) an early fast decaying pattern; 3) a slowly increasing pattern; and 4) an oscillating pattern (Figure 3A); i.e., the pattern of canonical trajectories (VL, D, I, O) in Table 1. In contrast to the observed iPDP similarities, we find that worm and fly have very different external principal dynamic patterns (ePDPs) (Figure 3B); i.e., the canonical temporal expression trajectories driven by species-specific TFs. The meta-gene canonical temporal expression trajectories driven by the worm-specific regulatory network; i.e., worm ePDPs, include a varied pattern at late embryonic development stage, a varied pattern that occurs early during the embryonic development, a fast increasing and then unvarying pattern, a decaying pattern, and an increasing pattern late during the embryonic development. The fly ePDPs, however, have two fast decaying patterns early during the embryonic development, a fast increasing pattern at a later stage during the embryonic development, and a highly increasing oscillation pattern. In addition, we checked the sensitivity of iPDP/ePDPs to small perturbations to internal/external regulatory networks by the leave-one-out method; i.e., we removed one gene in the internal/external group, ran DREISS, and obtained the ordered iPDP/ePDP eigenvalues for the remaining genes. We repeated the leave-one-out method for all genes, and finally found the ranges in which iPDP/ePDP eigenvalues vary shown as error bars in Figure S1. We can see that the iPDP eigenvalues vary less than ePDP ones for both worm and fly, which implies that the principal dynamic patterns of worm-fly orthologous genes driven by their conserved regulatory network are more robust to small changes than ones driven by their species-specific regulatory networks.

The above results suggest that the conserved regulatory networks from orthologous meta-genes between worm and fly have similar effects to orthologous meta-genes, given their similar iPDPs (i.e., both have four patterns, as described above). The species-specific regulatory networks from species-specific meta-genes (i.e., worm-specific or fly specific TFs) have effects that differ from the orthologous meta-genes for their different ePDPs.

## Orthologous genes have correlated coefficients between worm and fly for their matched internal principal dynamic patterns

In both worm and fly, we observe the similar four types of internally driven canonical temporal expression trajectories; i.e., internal principal dynamic patterns (iPDPs). Thus, we are interested in seeing how individual orthologous genes relate to those dynamic patterns. We find that the worm-fly orthologous genes have correlated coefficients over each of the four iPDPs. Based on Equation (10), we can obtain the coefficients of orthologous genes for each iPDP. We find that their coefficients are significantly correlated between worm and fly iPDPs with a similar pattern (See patterns in Table 1): *r*=0.33 (*p*<2.2e-16) for the highly varied pattern at late embryonic development stages, *r*=0.66 (*p*<2.2e-16) for the fast decaying pattern at early embryonic development stages, *r*=0.67 (*p*<2.2e-16) for the slowly increasing pattern during embryonic development, and *r*=0.73 (*p*<2.2e-16) for the oscillation pattern during embryonic development (Figure 4), where *r* represents Spearman correlation of iPDP coefficients of orthologous genes between worm and fly. This implies that, not only do the orthologous meta-genes have similar internal (conserved) regulatory effects (i.e., similar iPDPs), but the worm-fly orthologous genes also have similar internally-driven expression dynamics as resulted from their significantly correlated coefficients for iPDPs. The ePDPs between worm and fly generally do not show a high degree of matching similarity, but if we flip worm ePDP No. 3, and compare with fly ePDPs No. 4 and No. 5, all three of them are roughly representing the fast decaying patterns. We find that orthologous gene correlation coefficients between the ePDP patterns are very small (Spearman correlation *r*=0.12 of the orthologous gene coefficients of worm ePDP No.3 and fly ePDP No. 4, and *r*=0.18 of worm ePDP No. 3 vs. fly ePDP No. 5).

## Ribosomal genes have significantly larger coefficients for the internal than external principal dynamic patterns, but signaling genes exhibit the opposite trend

The ribosome produces proteins, which is an ancient process and conserved across worm and fly, organisms separated by almost a billion years of evolution. The ribosomal genes are highly expressed during embryogenesis, since intensive cell division and migration require a large amount of proteins to be synthesized. We collected 195 ribosome-related genes based on the GO annotations. We compared the iPDP and ePDP coefficients of ribosomal genes, and found that the iPDP coefficients are significantly larger than ePDP ones in both worm (KS-test *p*<0.001) and fly (KS-test *p*<2.2e-16) as shown in Figure 5A. This means that the ribosomal gene expression is significantly more influenced by the conserved regulatory network than by the species-specific regulatory network, which is consistent with ribosomal genes having conserved functions during embryonic development.

The orthologous genes related to signal transduction for cell-cell communication (a significantly more recent evolutionary adaptation relative to the ribosome) exhibit the opposite trend. We found that 320 signaling genes from GO annotations have significantly larger ePDP coefficients than iPDP ones in both worm (KS-test *p*<7e-4) and fly (KS-test *p*<6e-4), as shown in Figure 5B. This result implies that the signaling gene expression is significantly more driven by the species-specific regulatory network than by the conserved regulatory network, which is consistent with the signaling genes being commonly associated with species-specific functions, such as body plan establishment and cell differentiation.

## DNA replication and Proteasome machinery are enriched in orthologous genes with high coefficients for the dynamic patterns with fast growing canonical trajectories

We next turn to the biological meaning of individual canonical temporal expression trajectory for iPDPs and ePDPs. For the fast-decaying pattern (2nd iPDP), we find that the DNA replication is significantly enriched in Top 300 (~10%) orthologous genes that have the most negative coefficients for this pattern, in both worm (*p*<1.6e-8) and fly (*p*<4.5e-6). The GO enrichment analysis was performed using DAVID [[23](#_ENREF_23)]. The very negative coefficients for the fast decaying pattern mean high positive coefficients for a fast-growing pattern (vertically flipped 2nd iPDP), showing a drastic increase at the beginning of embryogenesis, then remain flat during the late embryogenesis (red curves in Figure 6). Most of the cell division of embryogenesis in both worm and fly happens approximately within the first 300 minutes. Then, the cell elongation and migration start to dominate the development [[24](#_ENREF_24),[25](#_ENREF_25)]. The mRNA abundance of the genes involved in DNA replication may change accordingly. This is well reflected by the second iPDP. Interestingly, the original expression patterns of those top orthologous genes actually do not have fast-growing patterns (black curves in Figure 6), probably because of the combined effects of both conserved and species-specific GRN. Maternal mRNAs, which are pre-loaded before fertilization, may also mask the fast growing pattern of DNA replication genes. This pattern could only be observed after we separated the effect of two types of TFs using DREISS. In addtion, we did not find any enrichment of DNA replication in top genes of other iPDPs and ePDPs (*p*>0.05). Therefore, the iPDP patterns identified by our method reveal elementary cellular process of both species (i.e. DNA replication), which should mainly be controlled by the conserved regulatory network.

Besides a fast growing pattern driven by conserved TFs, we also identified a fast growing pattern driven by non-conserved TFs for the two species. The Top 300 orthologous genes (~10%) with fast-growing worm and fly ePDPs (i.e., driven by species-specific regulatory networks) shared 36 orthologous genes. 10 of them encode proteins in the proteasome complex (*p*<1.2e-9). Protein degradation is not only a key process in apoptosis, but also throughout the entire course of development [[26](#_ENREF_26),[27](#_ENREF_27)]. For example, eliminating proteins that are no longer needed is a vital process during embryo development; e.g., the maternal proteins need to be cleaned as the embryogenesis proceeds). Previous reports also showed that different species usually have different maternal mRNA in the oocyte, which indicates that species-specific strategies might be utilized to regulate the protein degradation process [[28](#_ENREF_28)]. In this study, after separating the effect of conserved and non-conserved regulatory networks, we observed that the protein degradation is significantly enriched in the genes majorly driven by species-specific TFs.

Besides the 36 shared genes in the fast-growing pattern driven by species-specific TFs, we also noted a couple of interesting observations. Among the Top 300 worm orthologous genes with fast-growing ePDPs, genes involved in calcium ion binding (*p*<2e-6), GTP binding (*p*<7e-3) and neuron differentiation (*p*<0.05) are over-represented, suggesting that they are activated in the early stage of embryogenesis by worm-specific TFs. This observation indicates the GRN of these genes have evolved after the speciation. Proteins involved in calcium ion binding or GTP binding usually play a role in cell signal transduction [[29](#_ENREF_29)]. In fact, the genes involved in Wnt signaling and MAPK signaling exhibits a two-fold change.

In contrast, the Top 300 fly genes with a fast-growing ePDP show no enrichment in signaling transduction or cell differentiation. Instead, functions associated with respiration, such as oxidative phosphorylation, are enriched (*p*<5e-10). The enrichment of energy generation in the Top 300 fly genes with a fast-growing ePDP is probably indicative of the large energy requirement during fly embryogenesis [[30](#_ENREF_30)], which did not provide the evolutionary conservation of this energy-related gene regulation. Our result reveals that the fly genes associated with respiration are more up-regulated by fly-specific TFs relative to conserved TFs, and that this up-regulation evolved after the separation of worm and fly. In addition, the lack of signaling enrichment might be due to the different sampling time points. It is well-known that the Wnt signaling in worms starts as early as at the 4-cell stage, when one cell receives the signal and starts differentiation [[31](#_ENREF_31)]. The time-series worm transcriptome data used in our study may have the resolution to detect those processes. However, since each of the first 10 cell cycles takes less than 10 minutes in the fly embryo [[32](#_ENREF_32)], the 2 hour time interval in fly data may not have the resolution to capture the early regulatory events, such as Wnt signaling.

# Discussion

In this paper, we presented a novel computational method, DREISS, which decomposes time-series expression data of a group of genes into the components driven by the regulatory network inside the group (internal regulatory subsystem), and the components driven by the external regulatory network consisting of regulators outside the group (external regulatory subsystem). DREISS is a general-purpose tool that can be used to study the gene regulatory effects of any interested biological subsystems such as protein-coding transcription factors, micro-RNAs, epigenetic factors and so on. As an illustration, we applied DREISS to the time-series gene expression datasets for worm and fly embryonic developments from the modENCODE project [[10](#_ENREF_10)], and compared the worm-fly orthologous gene expression dynamic patterns driven by the conserved regulatory network (i.e., regulation effects from orthologous TFs), with the patterns driven by the species-specific regulatory networks (i.e., regulation effects from worm or fly specific TFs). We found that the conserved TFs drive similar genomic functions, but non-conserved TFs drive species-specific functions of orthologous genes between worm and fly, implying that, in addition to having ancient conserved functions, orthologous genes have been regulated by evolutionarily younger GRNs to execute species-specific functions during the evolution. This work can be easily extended to study the regulatory effects from orthologous TFs and species-specific TFs to species-specific genes. For example, one can find the expression dynamic patterns of worm/fly specific genes driven by specific TFs, and identify the genes with strong patterns associated with worm/fly specific functions, such as body formations. To the best of our knowledge, DREISS is the first method to reveal how the evolution of GRNs affects gene expression during embryogenesis.

We emphasize that DREISS is a general-purpose method (a free downloadable R tool available from github.com/gersteinlab/dreiss). Users can define the internal group (*X*) and external group (*U*) according to their interests. For example, if users want to identify the protein-coding expression patterns driven by miRNAs, they can define miRNAs as an external group and protein-coding genes as an internal group. Additionally, DREISS can be applied to more than two datasets, such as comparing worm, fly and human embryonic stem cell developmental data, and finding their conserved and specific developmental expression patterns. The expression patterns driven by human-specific regulatory factors will potentially help us understand human-specific developmental processes along with the associated human genes.

Due to the limited time samples in gene expression datasets, DREISS uses the simple linear state space model (i.e. the first order linear invariant difference equation) to model the temporal gene expression dynamics, and identify principal temporal dynamic patterns. This model assumes that the gene regulatory networks controlling temporal gene expression dynamics does not change across the entire biological process such as (*A*, *B*) in Equation (1). Thus, based on the analytic analysis, the principal dynamic patterns (PDPs) must follow a small set of canonical temporal trajectories (Table 1). With the rapidly increasing gene expression data, we can extend DREISS to more advanced models such as switched and hybrid system models, non-linear models [[33](#_ENREF_33)], which will allow us to study the gene regulatory networks are time varying, and potentially find the more temporal gene expression patterns capturing the more complex gene regulatory activities.

Figure captions

**Figure 1 DREISS workflow**. **1:** DREISS models temporal gene expression dynamics using state-space models in control theory. The “state” refers to the expressions for a large group of genes of interest, such as the worm-fly orthologous genes investigated here. The “control” refers to any other group of genes that contribute to gene expressions of the “state”, such as the species-specific TF studied here. **2**: it then projects high-dimensional gene expression space to lower-dimensional meta-gene expression spaces using dimensionality reduction techniques.

**3**: it derives the effective state-space models for meta-genes so that model parameters can be estimated. **4**: it then identifies the meta-gene expression dynamic patterns; i.e., canonical temporal expression trajectories driven by “state” (internal) and by “control” (external) based on the analytic solutions to estimated models. **5**: it finally calculates the coefficients of genes for the dynamic patterns of linear transformations between genes and meta-genes.

**Figure 2 State space model for genes and the effective model for meta-genes. A)** linear state space model for a given subsystem’s gene expression; i.e., linear first-order difference equations in Equation (2), is used to formulate temporal gene expression dynamics for a given subsystem, the gene group *X* (comprising *N*1 genes) with external regulations from the gene group *U* (comprising *N2* genes) at time points 1, 2, … , *T*. The vector  , the “state”, includes *N*1 gene expression levels at time *t* in group *X*, and the vector  , the “input or control”, includes *N*2 gene expression levels at time *t* in group *U*. The system matrix  captures internal causal interactions among genes in *X* (i.e., the *i*th, *j*th element of *A*, *Aij* describes the contribution from the *j*th gene expression at time *t* to the *i*th gene expression at the next time *t*+1). The control matrix  captures external causal regulations from the genes in *U* to genes in *X* (i.e., the *i*th, *j*th element of *B*, *Bij* describes the contribution from the *j*th gene expression in *U* at time *t* to the *i*th gene expression in *X* at the next time *t*+1). **B)** Meta-gene expression levels. The meta-gene expression levels are obtained by , where , the “meta-gene state”, includes *M*1 (<< *N*1 and <*T*) meta-gene expression levels; i.e., the first *M*1 elements of the *t*th row of the matrix whose columns are right-singular vectors of the matrix  in group *X* by the singular value decomposition (SVD) [[19](#_ENREF_19)]; the vector  , the “meta-gene input or control”, includes *M*2 (<< *N*2 and <*T*)meta-gene expression levels (i.e., the first *M*2 elements of the *t*th row of the matrix whose columns are right-singular vectors of the matrix SVD of matrix  at time *t* in group *U*; is the linear projection matrix of SVD from *M*1 meta-gene expression space to *N*1 gene expression space in *X,*  is the linear projection matrix of SVD from *M*2 meta-gene expression space to *N*2 gene expression space in *U)*, and (.)\* is a pseudo-inverse operation; i.e., *W\*W*=*I*, where *I* is the identity matrix. **C)** Effective state space model for meta-genes. The effective state-space model for meta-genes, Equation (4) is obtained by using linear projections *WX* and *WU* between genes and meta-genes from Equations (1-3). The effective meta-gene system matrix  captures internal causal interactions among meta-genes in *X* (i.e., the *i*th, *j*th element of  () describes the contribution from the *j*th meta-gene expression at time *t* to *i*th meta-gene expression at next time *t*+1), and the effective control matrix  captures external causal regulations from meta-genes in *U* to meta-genes in *X* (i.e., the *i*th, *j*th element of , describes the contribution from the *j*th meta-gene expression in *U* at time *t* to *i*th meta-gene expression in *X* at next time *t*+1). Equation (4) describes the effective state space model for the meta-genes in X, whose expression dynamics are determined byand. Because the meta-gene dimension, *M*1 (*M*2) is less than *T*, and much less than *N*1 (*N*2), we can estimateand as follows.

**Figure 3 Principal dynamic patterns of orthologous genes between worm and fly during embryonic development. A)** Metagenes of orthologous genes have similar internal driven principal dynamic patterns.Meta-gene canonical temporal expression trajectories driven by conserved regulatory networks (i.e., internal principal dynamic patterns, iPDPs) include four major patterns in both worm and fly embryonic development: 1) a highly varied pattern late (iPDP with the real eigenvalue No. 1); 2) a fast decaying pattern early (iPDP with the real eigenvalue No. 2); 3) a slowly increasing pattern (iPDP with the real eigenvalue No. 3); and 4) an oscillating pattern (iPDP with the complex eigenvalue). **B)** Metagenes of orthologous genes have different external driven principal dynamic patterns. Worm and fly have very different external principal dynamic patterns (ePDPs); i.e., the canonical temporal expression trajectories driven by species-specific TFs. The meta-gene dynamic patterns driven by the worm-specific regulatory network; i.e., worm ePDPs consist of a varied pattern at late embryonic development (real eigenvalue No. 1), a varied pattern at early embryonic development (real eigenvalue No. 2), a fast increasing and then unvarying pattern (real eigenvalue No. 3), a decaying pattern (real eigenvalue No. 4), and an increasing pattern at late embryonic development (real eigenvalue No. 5). The fly ePDPs, however, have two fast decaying patterns at early embryonic development (real eigenvalue No. 1 and 2), a fast increasing pattern at late embryonic development (real eigenvalue No. 3), and a highly increasing oscillation pattern (complex eigenvalue).

**Figure 4 Orthologous genes have correlated coefficients between worm and fly for their matched internal principal dynamic patterns**. The worm-fly orthologous genes have correlated coefficients over each of four iPDPs. Their coefficients are significantly correlated between worm and fly iPDPs with a similar pattern: r=0.33 (p<2.2e-16) for the highly varied pattern at late embryonic development, r=0.66 (p<2.2e-16) for the fast decaying pattern at early embryonic development, r=0.67 (p<2.2e-16) for the slowly increasing pattern during embryonic development, and r=0.73 (p<2.2e-16) for the oscillation pattern during embryonic development.

**Figure 5 Ribosomal genes have significantly larger coefficients for internal than external principal dynamic patterns, but signaling genes exhibit the opposite trend**. **A**) The iPDP and ePDP coefficients of ribosomal genes are compared: the iPDP coefficients are significantly larger than ePDP ones in both worm (KS-test p<0.001) and fly (KS-test p<2.2e-16); **B**) The iPDP and ePDP coefficients of signaling genes (cell-cell communication) are compared: they have significantly larger ePDP coefficients than iPDP ones in both worm (KS-test p<7e-4) and fly (KS-test p<6e-4).

**Figure 6 DNA replication is enriched in orthologous genes with high coefficients for the dynamic patterns with fast growing canonical trajectories**. For the fast-decaying pattern (2nd iPDP), we found that the DNA replication is significantly enriched in Top 300 (~10%) orthologous genes that have the most negative coefficients for this pattern, in both worm (p<1.6e-8) and fly (p<4.5e-6). The very negative coefficients for the fast decaying pattern means high positive coefficients for a fast-growing pattern, showing a drastic increase at the beginning of embryogenesis, then remain flat during the late embryogenesis (red curves). The original expression patterns of those top orthologous genes actually do not have fast-growing patterns (black curves).

**Figure S1 Principal dynamic patterns and their eigenvalues. A)** internal principal dynamic patterns (PDPs); **B)** external PDPs of orthologs during worm and fly embryonic development. Barplots show the eigenvalues of PDPs. The error bar for each eigenvalue tells the its variation range. We left one gene out, and calculated eigenvalues for the remaining genes thus obtaining the eigenvalue variations. The curves show the canonical temporal expression trajectories of PDPs.

References

1. Kim PM, Tidor B (2003) Subsystem identification through dimensionality reduction of large-scale gene expression data. Genome Res 13: 1706-1718.

2. Vilar JM (2006) Modularizing gene regulation. Mol Syst Biol 2: 2006 0016.

3. Peter IS, Davidson EH (2011) Evolution of gene regulatory networks controlling body plan development. Cell 144: 970-985.

4. Chen K, Rajewsky N (2007) The evolution of gene regulation by transcription factors and microRNAs. Nat Rev Genet 8: 93-103.

5. Levin M, Hashimshony T, Wagner F, Yanai I (2012) Developmental milestones punctuate gene expression in the Caenorhabditis embryo. Dev Cell 22: 1101-1108.

6. Kalinka AT, Varga KM, Gerrard DT, Preibisch S, Corcoran DL, et al. (2010) Gene expression divergence recapitulates the developmental hourglass model. Nature 468: 811-814.

7. Yanai I, Peshkin L, Jorgensen P, Kirschner MW (2011) Mapping gene expression in two Xenopus species: evolutionary constraints and developmental flexibility. Dev Cell 20: 483-496.

8. Irie N, Kuratani S (2011) Comparative transcriptome analysis reveals vertebrate phylotypic period during organogenesis. Nat Commun 2: 248.

9. Casci T (2011) Development: Hourglass theory gets molecular approval. Nat Rev Genet 12: 76.

10. Gerstein MB, Rozowsky J, Yan KK, Wang D, Cheng C, et al. (2014) Comparative analysis of the transcriptome across distant species. Nature 512: 445-448.

11. Brogan WL (1991) Modern control theory. Englewood Cliffs, N.J.: Prentice Hall. xviii, 653 p. p.

12. Rangel C, Angus J, Ghahramani Z, Lioumi M, Sotheran E, et al. (2004) Modeling T-cell activation using gene expression profiling and state-space models. Bioinformatics 20: 1361-1372.

13. Bansal M, Della Gatta G, di Bernardo D (2006) Inference of gene regulatory networks and compound mode of action from time course gene expression profiles. Bioinformatics 22: 815-822.

14. Huang S, Ingber DE (2006) A non-genetic basis for cancer progression and metastasis: self-organizing attractors in cell regulatory networks. Breast Dis 26: 27-54.

15. Saeys Y, Inza I, Larranaga P (2007) A review of feature selection techniques in bioinformatics. Bioinformatics 23: 2507-2517.

16. Chu S, DeRisi J, Eisen M, Mulholland J, Botstein D, et al. (1998) The transcriptional program of sporulation in budding yeast. Science 282: 699-705.

17. Wang D, Arapostathis A, Wilke CO, Markey MK (2012) Principal-oscillation-pattern analysis of gene expression. PLoS One 7: e28805.

18. Wang D, Markey MK, Wilke CO, Arapostathis A (2012) Eigen-genomic system dynamic-pattern analysis (ESDA): modeling mRNA degradation and self-regulation. IEEE/ACM Trans Comput Biol Bioinform 9: 430-437.

19. Golub GH, Van Loan CF (1996) Matrix computations. Baltimore: Johns Hopkins University Press. xxvii, 694 p. p.

20. Cull P, Flahive ME, Robson RO (2005) Difference equations : from rabbits to chaos. New York: Springer. xiii, 392 p. p.

21. Reece-Hoyes JS, Deplancke B, Shingles J, Grove CA, Hope IA, et al. (2005) A compendium of Caenorhabditis elegans regulatory transcription factors: a resource for mapping transcription regulatory networks. Genome Biol 6: R110.

22. Shazman S, Lee H, Socol Y, Mann RS, Honig B (2014) OnTheFly: a database of Drosophila melanogaster transcription factors and their binding sites. Nucleic Acids Res 42: D167-171.

23. Huang da W, Sherman BT, Lempicki RA (2009) Systematic and integrative analysis of large gene lists using DAVID bioinformatics resources. Nat Protoc 4: 44-57.

24. Bate M, Martinez Arias A (1993) The Development of Drosophila melanogaster. Plainview, N.Y.: Cold Spring Harbor Laboratory Press.

25. Baugh LR, Hill AA, Slonim DK, Brown EL, Hunter CP (2003) Composition and dynamics of the Caenorhabditis elegans early embryonic transcriptome. Development 130: 889-900.

26. DeRenzo C, Seydoux G (2004) A clean start: degradation of maternal proteins at the oocyte-to-embryo transition. Trends Cell Biol 14: 420-426.

27. Du Z, He F, Yu Z, Bowerman B, Bao Z (2015) E3 ubiquitin ligases promote progression of differentiation during C. elegans embryogenesis. Dev Biol 398: 267-279.

28. Shen-Orr SS, Pilpel Y, Hunter CP (2010) Composition and regulation of maternal and zygotic transcriptomes reflects species-specific reproductive mode. Genome Biol 11: R58.

29. Aspenstrom P (2004) Integration of signalling pathways regulated by small GTPases and calcium. Biochim Biophys Acta 1742: 51-58.

30. Tennessen JM, Bertagnolli NM, Evans J, Sieber MH, Cox J, et al. (2014) Coordinated metabolic transitions during Drosophila embryogenesis and the onset of aerobic glycolysis. G3 (Bethesda) 4: 839-850.

31. Sawa H, Korswagen HC (2013) Wnt signaling in C. elegans. WormBook: 1-30.

32. Gilbert SF (2000) Developmental biology. Sunderland, Mass.: Sinauer Associates. xviii, 749 p. p.

33. Schaft AJvd, Schumacher JM (2000) An introduction to hybrid dynamical systems. London ; New York: Springer. xi, 174 p. p.

1. [↑](#footnote-ref-2)