MODELING GENES INTERACTION: FITTING CHEMICAL KINETICS ORDINARY DIFFERENTIAL EQUATIONS TO MICROARRAY DATA

A Dissertation
Presented to
the Faculty of the Department of Mathematics
University of Houston

In Partial Fulfillment
of the Requirements for the Degree
Doctor of Philosophy

By
Zijun Luo
August 2010
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Zijun Luo

APPROVED:

Prof. Robert Azencott,
Chairman

Prof. Preethi Gunaratne

Prof. Kresimir Josic

Prof. Andrew Torok

Prof. Cecilia Williams

Dean, College of Natural Sciences and Mathematics
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Abstract

This thesis focuses on an active area in mathematics applied to biology, namely the modelling of genes interactions controlling the differentiation of embryonic stem cells, starting from MicroArray data. We have developed and implemented novel algorithmic approaches to this question, and tested them on two very large microarray data sets obtained by Austin Cooney’s Lab at Baylor College of Medicine.

A key biological question was to exploit these data to elucidate, in a set of over 20,000 messenger RNA (mRNA) genes, which mRNAs are actually repressed by each micro-RNA (miRNA) in a specific list of 266 known micro-RNAs. Recall here that microRNAs are short RNA strands which exert their inhibiting functions by binding their target mRNAs. Since the microarray data studied involve the simultaneous time evolution of expression levels for 20,000 mRNA genes and 266 micro-RNAs, identifying the specific genes interactions just mentioned is not an easy task. The algorithmic technique we have developed is to formalize potential genes interactions by explicit chemical kinetics equations (CKEs) parametrized by unknown parameters, and then to compute good estimates of these parameters.

An essential technical problem was to derive the adequate types of nonlinear ODEs (ordinary differential equations) to be used to model the chemically plausible CKEs, and to estimate the unknown parameters of these CKEs by algorithmic analysis of the time evolutions for expressions data.

One of the challenges is the massive size of the microarray data and the huge combinatorial possibilities number of groups of genes which may actually interact. Another statistical and mathematical challenge was to enforce a parameter parsimony principle, to avoid the massive and not very meaningful over-parameterization.
In particular the thesis has focused on the intensive modeling and validation (or invalidation) of more than 5,000 biologically plausible instances of two basic architectural motifs for miRNA interactions with the main genes controlling differentiation in ES cells. These basic motifs are of small size and always involve at least one pair of potentially interacting mRNA and miRNA, in systematic compliance with major biological reference tables listing potential (mRNA, miRNA) interactions.
## Contents

1 Introduction ........................................... 1

2 Biological Background ................................. 6
   2.1 Embryonic Stem Cells (ES-cells) ................. 6
   2.2 The Main Genes Controlling ES Cells Differentiation ... 7
   2.3 Microarray Data: Experimental Acquisition Modalities .... 8

3 A Specific Microarray Dataset for ES-cells Differentiation .... 10
   3.1 Detailed Description of ES-cells Microarray Data .......... 10
   3.2 Synthesis of Multi-Chips Recordings for MicroRNA Data . .... 12
   3.3 Data Interpolation by Piecewise Cubic Hermite polynomials .... 22
   3.4 Sensitivity of Data Interpolation to Measurements Errors ... 22
   3.5 Western Blot Data for 4 Key Proteins .................. 24

4 Classical Microarray Data Analysis ..................... 28
   4.1 Visualization of Microarray data ...................... 28
   4.2 Correlation Analysis of Microarray Data ................. 29
   4.3 Microarray Data Analysis: Clustering .................... 30
      4.3.1 Microarray Data Analysis: Principal Component Analysis (PCA) ... 32
   4.4 Microarray Data Analysis: Machine Learning Methods .......... 33
   4.5 Pretreatments to Adjust for Dye Differences ............... 33
5 Previous Results Linking miRNAs and Regulatory Loops for ES Cells Differentiation

5.1 Sequence Analysis and Reference Tables for Genes Interaction

5.2 Previous Microarray Data Analysis

5.3 Previous Biological Interpretation of correlation analysis

5.4 Open Questions Left Unresolved in Previous Study

6 Basic Architectural Motifs for Regulatory Loops

6.1 the Main Interaction Motifs to Model the Impact of MiRNAs on the Regulatory Loops of Differentiation

6.2 Motif A Architectures Linking miRMNAs and the ES Cells Regulatory Network

6.3 Key Families of Motif A Architectures

6.4 Motif B Architectures Linking miRMNAs and the ES Cells Regulatory Network

6.5 Modeling and Validation Methodology for Motif A and Motif B architectures

7 Modelling Basic Interaction Motifs by Chemical Kinetics Equations

7.1 Modeling Microarray Data by Chemical Kinetics Equation

7.2 Chemical Kinetics Equation for Motif A

7.2.1 Post-transcriptional Repressors and Architectural Interaction Motif A

7.3 Chemical Kinetics Equation for Translation Repressors

7.3.1 Interaction Motif B

7.3.2 CKEs to model motif B architectures

7.4 Biochemical Assumptions and CKEs Derivation for Motifs A and B

7.5 Simplified Chemical Kinetics Equations for High Degradation Rate

7.6 Formal Invariance of our CKE models by Scale Changes

8 Parameter Estimation for the CKE models of Motifs A and B

8.1 Parameter Estimation for Nonlinear CKEs

8.1.1 Fitting Chemical Kinetics Dynamic Systems to Data
Chapter 1

Introduction

Transcriptional and translational regulations are two fundamental mechanisms of living cells. These two processes always involve interactions of three main molecular species: messenger RNA (mRNA), proteins associated to the mRNAs, and micro-RNAs (miRNA). One of the challenges in bioinformatics is to analyze recorded genes expression levels (and/or genes structure) to determine how key genes are regulated and with which other genes or associated proteins they interact within circumscribed molecular networks.

Transcription is the first step leading to gene expression. The stretch of DNA transcribed into an RNA molecule is called a transcription unit and encodes at least one gene. If the gene transcribed encodes for a protein $P$, the transcription result is a messenger RNA (mRNA) $G$. The mRNA $G$ transports the protein coding information to the sites of protein $P$ synthesis, and will then be used to produce the protein $P$ via the process of translation.

MicroRNAs (miRNAs) are small non-coding RNAs of roughly 22 nucleotides in length, which are able to bind with and inhibit protein coding mRNAs through complementary
base pairing. The minimum requirement for this interaction is that at least six consecutive nucleotides undergo base pairing to establish a miRNA-mRNA duplex. Thus a given miRNA can potentially bind and silence hundreds of mRNAs across a number of signaling pathways. These miRNAs are able to integrate multiple genes into biologically meaningful networks underlyi

MicroRNAs regulate gene expression post-transcriptionally through two distinct modalities: they may directly downregulate their specific target mRNAs, but mostly they inhibit the translation of their target mRNAs. If an miRNA denoted M binds to one of its mRNA target G with partial complementarity, then the translation process of G will be inhibited [66, 67]. If M binds to G with perfect or near-perfect complementarity, then G is cleaved resulting in its decay [64, 65, 68].

MiRNA genes expression is ultimately controlled by the same transcription factors that regulate protein coding genes. These transcription factors themselves are under the regulation of multiple miRNAs some of which they also activate or repress. Consequently miRNAs and mRNAs interact with each other in complex feed-forward and feed-back networks [61, 62].

Embryonic Stem Cells (ES-cells) for mice and humans are pluripotent cells which can replicate indefinitely, and then differentiate into many quite distinct types of cells. ES cell pluripotency is regulated both by extrinsic signaling pathways and by intrinsic gene networks controlling ES-cells differentiation. These regulatory mechanisms involve a network of transcription factors including the key genes GCNF, Oct4, Sox2, Nanog, Tbx3, Essrb, Klf4, cMyc, Eed, Ezh1, Ezh2 [2]. In ES-cells, key transcription factors such as Oct4, Sox2 and Nanog are closely linked with micro-RNAs, which are definitely quite enriched in ES-cells, for both mice and humans [2, 69].

Genome wide analysis using microarray data and sequencing technologies have significantly
expanded our knowledge of the complex regulatory networks underlying properties unique to ES cells.

Microarray is a technology used to obtain simultaneous genomic expression level profiles for a large number of genes. It generates the time indexed expression levels of the thousands of genes represented on the microarray. An microarray data file includes for each recorded gene $G$, and at multiple synchronous time points, the mean and standard deviation of the expression level of gene $G$.

We have focused our study on new algorithmics enabling the analysis of two large microarray data sets, recording the dynamic expression profiles of 20,000 mRNA genes and 266 miRNAs. These data were recorded on two types of mice ES-cells, during differentiation induced by retinoic acid. Expression levels for 4 key proteins were also recorded in the same contexts (laboratory of Dr. A. Cooney, Baylor College of Medicine). Classical microarray data analysis had been previously performed for these microarray data sets, implemented mostly by qualitative inference based on massive correlation analysis, and leading to the publication of interesting biological conclusions in [2]. In this paper, microarray data correlation analysis was combined with sequence analysis results, which classically identify similar regions on distinct DNA sequences, via popular biological reference tables TargetScan and miRanda, to predict potentially interacting miRNA-mRNA pairs in ES-cells differentiation control [1][2].

To go beyond the results of [2], we have attempted to formalize and optimally parametrize chemical kinetics equations (CKEs) underlying the two basic mechanisms through which microRNAs repress their target mRNAs, in the post-transcriptional process. We have thus formalized the two main repressive modalities of miRNAs by two types of interaction architectures (motifs A and B) between miRNA, mRNA, and associated proteins. Motif A
models transcriptional and post-transcriptional regulation, while motif B models translational regulation.

We have derived adequate Chemical Kinetics Equations to model the dynamics of motif A and motif B architectures, inspired by chemical kinetics models used in other contexts by [3, 7, 9, 10, 34, 35, 74–80]. Each such formal ODE is parametrized by a small number of unknown parameters (less than 10 parameters for each CKE). We have generated two lists (called ListA and ListB) of potential motifs to be evaluated, partially based on questions and hypotheses left open in [2]. To significantly narrow ListA, we have restricted the potentially interacting (miRNA,mRNA) pairs involved in each motif instance to be compatible with the predictions of the reference tables miRanda and TargetScan. For ListB, we have deliberately limited the number of miRNA repressors involved in each motif instance in order to keep a reasonably low value for the ratio of the number of parameters over the number of data points.

Mathematical modeling by nonlinear chemical kinetics equations generates complicated parameter estimation problems when one tries to adequately fit recorded microarray data. Generic cost function minimization techniques can be formally applied to parameter estimation, but they generally involve large amounts of computing time in our context, where we had to actually model and parametrize several thousands of architectural motifs, due to the the large number of potential genes interaction sub-architectures of interest in our context.

We have developed an innovative specific fast algorithms dedicated to parameter estimation for the two types of CKEs we have used. The algorithms we have implemented also provides relatively high-quality optimization for the quality of fit, since they integrate a blend of global search and local cost minimization. Algorithmic parametrization of the CKEs modeling each motif in ListA and ListB is then performed to optimize the fit of
these models with our two sets of WT and KO microarray data. We have also developed techniques to evaluate model robustness to the measurement errors corrupting the microarray data. We consider that a given interaction architecture of motif A or motif B type is “model validated” if our associated optimally parametrized model generates simulated predicted profiles close enough to the recorded expression level data. This methodology and the associated intensive computations we have performed thus generate several interesting families of “model validated” interacting miRNA-mRNA pairs involved in motif A architectures, as well as several families of model validated groups of miRNA repressors inhibiting specific mRNA generated proteins in motif B architectures. These model validated motif A or motif B architectures should be of interest to efficiently circumscribe further biological experiments, by focusing gene expression recordings on much smaller sets of miRNAs and mRNAs than those predicted by wide range reference tables miRanda or TargetScan.
Chapter 2

Biological Background

2.1 Embryonic Stem Cells (ES-cells)

Embryonic stem cells (ES cells) are pluripotent stem cells derived from the inner cell mass of the mammalian blastocyst.

Two key properties set ES cells apart from all the other cell types.

The first one is self-renewal, i.e. the ability to continuously replicate indefinitely as a result of their extensive proliferative potential.

The second is the property of pluripotency or the ability to develop into a number of different and specialized cells types through differentiation. In differentiation induced by Retinoic Acid (RA), ES cells begin in their undifferentiated state on Day 0, and following RA-induction start to differentiate on Day 1; the differentiation is complete by Day 6.

Genome wide analysis using microarray and sequencing technologies have significantly expanded our knowledge of the complex regulatory networks underlying properties unique to embryonic stem cells. We describe briefly these technologies in section [4] Data we model
and study here, two sets of mouse ES cells were treated by retinoic acid (RA) induction during differentiation, and time-course microarray data were recorded from day 0 to day 6.

The first set of cells includes only ES cells of wild type (WT), which refers to the ”normal” or ”standard” type of ES cells occurring in natural biological contexts. The second set of ES-cells consists of GCNF-KO cells, i.e. of mutant ES cells generated by chemically ”knocking out” the gene GCNF. These ES cells are engineered to carry only GCNF genes altered to become inoperative; altered GCNF genes will translate into nonfunctional proteins, if they are translated at all.

2.2 The Main Genes Controlling ES Cells Differentiation

ES cell pluripotence is regulated both by extrinsic signaling pathways and by intrinsic gene regulatory mechanisms [2] involving a network of transcription factors including GCNF, Oct4, Sox2, Nanog, Tbx3, Essrb, Klf4, cMyc, Eed, Ezh1, Ezh2 [2,20–32].

The orphan nuclear receptor GCNF protein (germ cell nuclear factor) is identified to be the transcriptional repressor of two key mRNA genes: Oct4 and Nanog [32]. The Oct4 and Nanog proteins are found to function together to regulate a significant proportion of their target genes (see [12]) in ES cells.

Oct4 and Nanog proteins influence the self renewal of ES cells by activating the self-renewal regulators (Sox2, Tbx3, Essrb, Klf4, cMyc), which maintains the self renewal process.

Oct4 and Nanog proteins influence ES pluripotency by activating the differentiation inhibitors (Ezh1, Ezh2, Eed), which suppress the differentiation process by repressing the Hox genes cluster. In animals, fungi and plants, the Hox genes are generally involved in
the regulation of patterns of development (morphogenesis). In ES cells, the Hox cluster activates the differentiation.

2.3 Microarray Data: Experimental Acquisition Modalities

A microarray is also called a DNA chip or a gene chip. It is the technology used to obtain simultaneous genomic profiles for a large number of genes (typically more than 10,000 genes).

The fundamental basis of DNA microarrays data acquisition is the process of hybridization. Two strands of nucleic acid, DNA or RNA, hybridize if they are complementary to each other. This principle is exploited to measure the unknown expression level of one RNA or DNA molecule (target) on the basis of the expression level measured for a complementary sequence (probe), that has hybridized with the target. The level of hybridization is usually quantified by optically measuring the level of a detectable chemical label, which "marks" or "tags" the target or the probe sequence in the experiment.

In the microarray technique, the probe sequences are immobilized on the bio-chip surface, and neighbouring probes are separated by a few micrometers only, so that one actually packs a very large number of distinct probes on a small single surface of 1 cm². Usually, the chemical labels become optically detectable through a fluorescent dye, which can be detected and quantified by a light scanner which analyzes the chip surface.

Each probe sequence matches a specific messenger RNA present in the sample. The concentration of a specific messenger RNA is a result from the expression level of its corresponding mRNA gene. At each acquisition time point, simultaneous optical scanning of all microarray spots records the simultaneous expression levels of all the target mRNAs present in the
sample; this generates the time indexed expression levels of all the genes represented on
the microarray. These recorded thousands of synchronous expression level profiles provide
then a quantitative simultaneous dynamic view of the time evolution for the underlying
biochemical process. The number of time points used to be quite small due to acquisition
costs, but this situation is quickly improving due to wider availability of cheaper acquisition
techniques.

Microarray after hybridization is scanned to be DAT file, which is the image of the scanned
array. Image analysis will then be performed and generates cell intensity files and chip de-
scription files. Processing software produces Excel file, CHP file or txt file, which are the
3 main formats for microarray data. An microarray data file stores the data information,
such as scanner, processing software, background level, back ground standard deviation,
probe-ID, average intensities, standard deviations and so fourth.
Chapter 3

A Specific Microarray Dataset for ES-cells Differentiation

3.1 Detailed Description of ES-cells Microarray Data

We have focused our study on a specific very large set of microarray data profiling the evolution profiles of 20,000 mRNA genes of mice ES cells, during differentiation induced by retinoic acid. These data have been presented and previously studied in [2] by quite classical correlation analysis techniques and visualizations of heat-maps displays. These microarray data for mRNAs have been acquired by the laboratory of Dr. Austin Cooney, Baylor College of Medicine. The other microarray data, focused on the recording of 266 known miRNA genes are provided by LC Science, inc.

The microarray data are in Excel format, each file contains 2 samples, one sample for standard wild type (WT) ES cells, and another one for GCNF-knock-out ES-cells. The miRNA data involve in particular 266 well identified micro-RNA genes, on which we
have focused our applicative study, in order to elucidate on which subgroups of the 20,000 recorded mRNA profiles these 266 miRNAs actually exert a repressive influence.

Data for 3 types of miRNA predictions, namely MCE-MIR (short for micro-conserved element miRNA prediction), Cand (short for candidates), MIR (short for miRNA prediction), and 266 identified miRNAs (mmu-mir, short for Mus musculus) are recorded. All microarray data are based on six probe replicates for each miRNA prediction (MCE-MIR, Cand, MIR) and eight probe replicates for miRNAs (mmu-mirs).

The miRNA data are recorded at days 0, 1, 3, 6 for both wild type (WT) and GCNF-knockout (KO).

The mRNA expression profiles, include the time evolution of expression levels for more than 20,000 mRNA genes, which involve 45,101 recordings since there are replicates.

These mRNA data are recorded on ES cells of WT type as well as on ES cells of GCNF-KO type, at days 0, 3, 6 using an Affymetrix mouse 430 2 array. Three biological replicates were performed per time point and thus 9 arrays were generated in total.

There are replicate arrays for miRNAs within each treatment. For WT and KO miRNAs data at day $t$, there are $K^t$ arrays (chips), each of which records signal intensities for the 266 miRNAs.

For day 0 of ES WT and KO data, there are 2 and 3 replicate arrays respectively.
For day 1 of ES WT and KO data, there are 2 and 1 replicate arrays respectively.
For day 3 of ES WT and KO data, there are 3 and 2 replicate arrays respectively.
For day 6 of ES WT and KO data, there are 2 and 2 replicate arrays respectively.

Recorded signal intensities are as usual assumed to be proportional to concentrations.

We naturally had to recombine redundant chips data as explained below.
3.2 Synthesis of Multi-Chips Recordings for MicroRNA Data

There are replicate miRNA arrays for both WT and KO ES-cells at each recording time point (day 0, 1, 3, 6) and we synthesize the replicate chips separately for WT and KO datasets. Denote by $M_{j,t}^k$ the recording of miRNA $j$ for day $t$ on chip $k$, where $k = 1, ..., K^t, t = 0, 1, 3, 6$, $j = 1, ..., 266$. For each pair $j, t$ we synthesize recordings by computing the average recording $avM_{j,t}$ over the 266 available recordings. Let

$$avM_{j,t} = 1/K^t \sum_k M_{j,t}^k$$

To synthesize the multiple recordings on mRNA replicates, we compute the averages across multiple recordings.

After synthesization of the data, we compute the mean value, variation, variation to mean, standard deviation and standard deviation to mean (also called dispersion ratio) of each observation for each gene (both miRNA and mRNA), and plotted the histograms for both WT and KO data. The mean values of miRNAs are distributed in higher values of intervals for WT than for KO (figure 3.2), which indicates that it is more likely that the miRNA expression is higher in WT than KO. The mean expression levels of mRNAs are distributed almost the same in both WT and KO context (figure 3.3). Variation is defined to be the difference of the maximum expression point and the minimum expression point for each observation. Figure 3.4 shows that the variations of miRNAs are a little bigger in WT than in KO, while figure 3.6 shows that the variation/mean ratio is bigger in KO than in WT. Figure 3.5 shows that the variations of mRNAs are bigger in WT than in KO, while figure 3.7 shows that the variation/mean ratio are also higher in WT than in KO. The dispersion ratios of miRNAs are lower in WT than in KO (figure 3.8), while the dispersion ratios of mRNAs are higher in WT than KO (figure 3.9).
Figure 3.1: Synthetic expression levels of 3 miRNAs of both WT and KO data.
Figure 3.2: Histograms of the mean values of the 266 miRNAs for both WT and KO data. The mean values are computed from 4 time-course data points of each miRNA.
Figure 3.3: Histograms of the mean values of the 45101 mRNAs for both WT and KO data. The mean values are computed from 3 time-course data points of each mRNA.
Figure 3.4: Histograms of the variations of the 266 miRNAs for both WT and KO data. The variation values are computed from the difference of the maximum measurement and minimum measurement for each miRNA.
Figure 3.5: Histograms of the variations of the 45101 mRNAs for both WT and KO data. The variation values are computed from the difference of the maximum measurement and minimum measurement for each mRNA.
Figure 3.6: Histograms of the variation/mean values of the 266 miRNAs for both WT and KO data. The values of variation is divided by mean value of each miRNA, and the lowest 3% mean values are taken out.
Figure 3.7: Histograms of the variation/mean values of the 45101 mRNAs for both WT and KO data. The values of variation is divided by mean value of each mRNA, and the lowest 3% mean values are taken out.
Figure 3.8: Histograms of the dispersion ratios of the 266 miRNAs for both WT and KO data. The values of standard deviation is divided by mean value of each miRNA, and the lowest 3% mean values are taken out.
Figure 3.9: Histograms of the dispersion ratios values of the 45101 mRNAs for both WT and KO data. The values of standard deviation is divided by mean value of each mRNA, and the lowest 3% mean values are taken out.
3.3 Data Interpolation by Piecewise Cubic Hermite polynomials

For each miRNA, and for each mRNA, we generate 19 interpolated concentration values at the 19 time points (\( t = 0, 1/3, 2/3, \ldots, 17/3, 18/3 = 6 \)) by Piecewise Cubic Hermite Interpolation (PCHIP) [48]. For example we interpolate a curve on \( n \) points \( f(x_k) = y_k, k = 1, \ldots, n \). On each subinterval \( x_k \leq x \leq x_{k+1} \), \( P(x) \) is the cubic Hermite interpolant to the given values and certain slopes at the two endpoints, with first derivative \( P'(x) \) is continuous but second derivative \( P''(x) \) is probably not continuous at \( x_k \). The slopes at the \( x_k \) are chosen to preserve the shape of the data and monotonicity. This means that, on intervals where the data are monotonic, so is \( P(x) \); at points where the data has a local extremum, so does \( P(x) \).

As shown in [48], such an interpolant may be more reasonable than a cubic spline if the data contains both “steep” and “flat” sections for this interpolation method preserves monotonicity and the basic qualitative features of concentrations dynamics.

3.4 Sensitivity of Data Interpolation to Measurements Errors

The evolutionary curves of the genes could vary after interpolation because of the measurement errors. For each expression value \( r(t) \) at time \( t \), \( t = 0, 3, 6 \) for mRNAs or \( t = 0, 1, 3, 6 \) for miRNAs, there is a corresponding measured standard deviation value \( \sigma(t) \). For the protein data we can take 5% as the relative error of the numerical data. We simulate 20 evolutions of each gene/protein following a uniform distribution with the measured or
Figure 3.10: **An example of interpolated evolution by PCHIP of miRNA mmu-let-7a in ES WT cells.** Blue solid line with circles on time 0, 1, 3, 6 is measured evolution. Red dash line is the interpolated curve.

artificial error at each recorded time point, and then interpolate these 20 evolutions. The measurements errors could sometimes be big, especially in the GCNF-knock-out data for the mRNAs (figure 3.11), so that the newly interpolated evolution curve could be very different in shape from the original curve. For miRNAs the simulated evolutions do not vary much from the measured evolution in most cases (figure 3.12), but there are large relative errors when the measured expressions are very low (figure 3.13). Since we set a low relative error 5% for proteins, figure 3.14 shows that the simulated evolutions are close to the measured evolution.
Figure 3.11: Simulate evolutions of mRNA Oct4 for WT (left) and KO (right). Blue solid line = interpolated evolution from the measured expression, green dash lines = interpolated evolutions from the simulated data.

3.5 Western Blot Data for 4 Key Proteins

For both WT and KO ES cells differentiation, protein expression levels have been recorded by Western Blots techniques implemented by Xueping Xu in Austin J. Cooney’s Laboratory (Baylor College of Medicine, Houston). These recordings were focused on the 4 proteins respectively associated to the genes GCNF, Oct4, Nanog, and Sox2. These genes are known to play important regulatory roles in ES-cells differentiation.

The Western blots recordings provide 4 data points for WT ES cells as well as for GCNF-KO ES cells, at time points (0, 1.5, 3, 6).
Figure 3.12: **Simulate evolutions of miRNA mmu-mir-296** for WT (left) and KO (right). Blue solid line=interpolated evolution from the measured expression, green dash lines=interpolated evolutions from the simulated data.

We have transformed the raw image data provided by Western Blots into digitized numerical grey scale values, by classical image analysis software tools [6]. After conversion of each blot into numerical image intensities, we have normalized the image intensities by the corresponding ACTIN intensities (which plays the part of an internal control). Then, as above, we have interpolated these normalized intensity data into 19 points at time points $t = 0, 1/3, ..., 18/3$ for both sets of ES-data (WT and GCNF-KO). Thus we obtained the evolutionary curves as shown in figure 3.16. Since the GCNF protein is knocked out in KO data, we simply set the expression level of GCNF at zero for KO data. These proteins data are assumed to be proportional to the protein concentrations.
Figure 3.13: **Simulate evolutions of miRNA mmu-mir-338 for WT (left) and KO (right).** Blue solid line=interpolated evolution from the measured expression, green dash lines=interpolated evolutions from the simulated data.

Figure 3.14: **Simulate evolutions of protein Nanog for WT (left) and KO (right).** Blue solid line=interpolated evolution from the measured expression, green dash lines=interpolated evolutions from the simulated data.
Figure 3.15: Western blots for 4 proteins and actins.

<table>
<thead>
<tr>
<th>Protein</th>
<th>WT</th>
<th>GCNF-/</th>
<th>RA(days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oct4</td>
<td></td>
<td></td>
<td>0 1.5 3 6</td>
</tr>
<tr>
<td>GCNF</td>
<td></td>
<td></td>
<td>0 1.5 3 6</td>
</tr>
<tr>
<td>Nanog</td>
<td></td>
<td></td>
<td>0 1.5 3 6</td>
</tr>
<tr>
<td>Actin</td>
<td></td>
<td></td>
<td>0 1.5 3 6</td>
</tr>
<tr>
<td>Sox2</td>
<td></td>
<td></td>
<td>0 1.5 3 6</td>
</tr>
</tbody>
</table>

Figure 3.16: Normalized expression levels of transformed protein data.
Chapter 4

Classical Microarray Data Analysis

4.1 Visualization of Microarray data

"Heat map" is a frequent graphic display modality used to visualize microarray data in terms of "log median ratios", which will be defined in the following paragraph.

Consider a sample of size $n$ of microarray data, which record expression levels (also called intensities) $g_i(t)$ for gene $G_i$ at time $t$, for $i = 1, \ldots, n$. For each $i$, let $\bar{g}_i$ be the median over time $t$ of the expression levels $g_i(t)$. Then for the $i^{th}$ gene, the log median ratio is defined by $\log_2(g_i(t)/\bar{g}_i)$. These ratios are classically displayed graphically as so-called Heat Maps. In the heat map [1,2] displayed below, red and green color squares respectively represent high and low values of the log median ratios, plotted as functions of time. We will see more examples of this type of visualization in section 5.2.
Figure 4.1: Example of Heat Map Visualization.

4.2 Correlation Analysis of Microarray Data

Established current techniques, like Sequence Analysis and Microarray Data Analysis provide essentially qualitative results about the interactions between the expression levels of the molecules of interest.

If we fix an mRNA gene denoted $G$, Sequence Analysis provides a long list denoted \( \text{List}(G) \) of potential miRNAs that may target $G$. Such lists are for instance available in tables such as miranda and Targetscan. Then extensive experiments would be required to tell which miRNAs in \( \text{List}(G) \) are actually targeting $G$, or to narrow the pool of miRNAs potentially targeting $G$.

Established microarray data analysis usually generates, for several thousands of genes $G_i$, the correlations between the expression levels of arbitrary pairs of genes, in order to identify the pairs $(G_i, G_j)$ having strongly positive or strongly negative correlation. This correlation analysis provides qualitative information about the potential direct interactions between the expression level of an miRNA and its potential mRNA targets. This qualitative analysis may also only detect indirect interactions between miRNAs and mRNAs,
since when an miRNA denoted \( M \) has strong negative correlation with an mRNA gene \( G \), it is quite possible that \( M \) does not target \( G \), but does repress the expression level of a protein \( P \) activating \( G \).

Therefore, this type of qualitative correlation analysis may generate a pool of potential miRNA candidates which seem to target a specific mRNA, but cannot generate a precisely descriptive model of the miRNA-mRNA interactions. Moreover, in order to quantitatively study the impacts of miRNAs in ES cells, one also has to take into account the expression level of proteins and model the interactions between miRNAs, their target mRNAs and the proteins associated to these targets.

We outline several algorithmic techniques which are very commonly used to identify smaller sets of predictive genes or pathways that could have closely related biological functions \[34, 46, 47\].

4.3 Microarray Data Analysis: Clustering

Cluster Analysis is used for grouping or segmenting a collection of vector valued observations into subsets or “clusters”, such that within each cluster, arbitrary pairs of observations are much more similar than pairs of observations located in distinct clusters. Central to the goals of cluster analysis is the notion of degree of similarity (or dissimilarity) between arbitrary pairs of vector observations being clustered.

Hierarchical Clustering methods require to quantify dissimilarity between disjoint groups of observations, based on user specified pairwise dissimilarities \( D(X_i, X_j) \) between observations \( X_i, X_j \). This method produces hierarchical representations in which the clusters at each level of the hierarchy are created by merging clusters at the next lower level. At the
lowest level, each cluster contains only one observation, while at the highest level there is only one cluster containing all observations. Two main strategies are used to implement hierarchical clustering.

The bottom-up strategies begin by associating a singleton cluster to each observation, and then recursively merge a selected pair of clusters into a single cluster.

Top-down strategies start with a single large cluster including all observations and recursively split one of the existing clusters into two new clusters. The cluster split at each iteration is often implemented by K-means, with \( K = 2 \), as in [54].

Another splitting approach (see [58]) defines dissimilarity \( D(X, C) \) between an observation \( X \) and any set \( C \) of observations as the average of the \( D(X, X') \) over all \( X' \) in \( C \). To split a cluster \( C \), start with the split \( H_0 = \text{"empty set"} \) and \( C_0 = C \); then generate iteratively new splits \( H_i, C_i \) of \( C \) by adding to \( H_i \) and eliminating from \( C_i \) one observation \( X \) selected in \( C_i \) in order to maximize \( D(X, C_i) - D(X, H_i) \). Stop the iterations when this maximal difference is negative.

K-means Clustering defines dissimilarity between vector observations \( X, X' \) by their squared Euclidean distance \( D(X, X') = ||X - X'||^2 \). One imposes a maximal number \( K \) of clusters \( C_1, \ldots, C_K \). Each observation \( X_i \) is assigned to only one cluster \( C_{g(i)} \). A cost function \( \text{Cost}(g) \) quantifies the clustering quality of the function \( g \) by the Total Cluster Variance

\[
TCV(g) = \frac{1}{2} \sum_{k=1}^{K} \sum_{g(i)=k} \sum_{g(j)=k} D(X_i, X_j)
\]

The K-means clustering algorithm minimizes \( TCV(g) \) by the following steps (see [54]) .

1. Call \( m_k \) the barycenter of the current cluster \( C_k \)

2. Define a new function \( \hat{g} \) by re-assigning each observation \( X_i \) to the index \( k = \hat{g}(i) \) minimizing \( D(X_i, m_k) \). This defines new clusters.
3. Iterate Steps 1 and 2 until the clusters stabilize.

4.3.1 Microarray Data Analysis: Principal Component Analysis (PCA)

A series of microarray experiments produces observations of differential expression for thousands of genes across multiple conditions. One problem is that different experiments seem different because of their biological context, but they may actually be identical or very similar in terms of relative genes expressions levels. In particular, correlation analysis may associate too tightly specific groups of genes, due to high redundancy for specific groups of measurements.

Principal Component Analysis (PCA) reduces the vector dimension of the data, and can help to separate the independent information contents of distinct experiments [55]. PCA classically extracts a small number of linear combinations of the observed vector variables, called principal components, and these principal components viewed as new observables will account for most of the variance in the observed variables. The principal components may then be used as predictor or criterion variables in subsequent analyses.

PCA involves the calculation of the eigenvalue decomposition for the data covariance matrix or singular value decomposition of a data matrix, usually after mean centering the data for each attribute [55]. Practical implementation may be performed via the MATLAB function "princomp".

PCA can be thought of as revealing the internal structure of the data in a way which best explains the variance in the data. If a multivariate dataset is visualized as a set of coordinates in a high-dimensional data space (1 axis per variable), PCA supplies the user with a lower-dimensional picture, a "shadow" of this object when viewed from its (in some sense) most informative viewpoint.
4.4 Microarray Data Analysis: Machine Learning Methods

For the analysis of microarray data, one of the ultimate goals is to estimate unknown dynamic relationships linking the expression levels of strongly interacting groups of genes. This broad formulation of course lends itself, at least formally, to multiple approaches by pure machine learning, with no biochemical modeling at all. We refer to [56] for a detailed survey of these approaches, which include for instance

- Multi-Layer Perceptrons (see [57,70])
- Support Vector Machines (SVM) (see [57,71,80])
- Self-Organizing Maps (SOM) (see [57])

We did not try to apply these techniques to the study of our data set, since we have privileged detailed modeling by optimal parametrizing of plausible chemical kinetics equations.

4.5 Pretreatments to Adjust for Dye Differences

Each miRNA chip used in this study was in digitized format and contained 2 samples of data (dyed in different colors), each of which included for each miRNA $M_i$, $i = 1, \ldots, 266$, both the measured expression level $m_i$ and the standard deviation $std_i$ of the corresponding error of measurement. Each one of these 2 files included also a measured background level $\mu$ and an estimated background standard deviation $\sigma$.

In each one of the two samples recorded by an miRNA chip, the expression levels $m_i$ were first centered by subtracting the background level $\mu$. Then these centered two-color data were normalized by the classical LOWESS filter.
LOWESS normalization, or Locally-Weighted Regression, is a technique for fitting a smoothing curve to a given dataset. LOWESS Intensity Dependent normalization is used for two-color data. It is a type of within-array normalization scheme which adjusts for intensity-dependent variation due to distinct dye properties. Dye bias is caused by inconsistencies in the relative fluorescence intensity between dyes Cy5 and Cy3 [44, 52]. These inconsistencies often result in nonlinear relationship between the fluorescences generated by these two dyes. Normalization is done separately for each array.

Denote the red and green intensities by $R_i$ and $G_i$ for $i = 1, \ldots, n = 266$. The adjusted ratio $r_i$ is computed by:

$$\log_2(r_i) = L_i \times \log_2(R_i/G_i)$$

where $L_i = L(\log_2(\sqrt{R_i \cdot G_i}))$ and $y = L(x)$ is the function $L$ generated by LOWESS fitting of the the $y_i = \log_2(R_i/G_i)$ to the $x_i = \log_2(\sqrt{R_i \cdot G_i})$. Then the adjusted red data $\hat{R}_i$ and green data $\hat{G}_i$ are given by

$$\log_2 \hat{G}_i = \log_2 G_i + \log_2 r_i$$

$$\log_2 \hat{R}_i = \log_2 R_i - \log_2 r_i$$

By applying LOWESS normalization to the data, the relationship between the log data of the two dyes becomes essentially linear.

Another approach to normalize two-color data is the classical quantile equalization, which is a technique to match two distinct probability distributions by nonlinear transformation of data generated by the second distribution. We refer for instance to [53] for a practical algorithm implementing quantile equallization. This approach is quite useful when the 2 distributions are fairly similar.
Chapter 5

Previous Results Linking miRNAs and Regulatory Loops for ES Cells Differentiation

Several publications indicate that miRNAs have important functions of post-transcriptional silencing and are involved in the regulation of differentiation in stem cells.

5.1 Sequence Analysis and Reference Tables for Genes Interaction

In bioinformatics, a sequence alignment is a way of arranging the sequences of DNA, RNA, or protein to identify regions of similarity that may be a consequence of functional, structural, or evolutionary relationships between the sequences [49]. Pairwise sequence
alignment methods are used to find the best-matching piecewise alignments of two query sequences. Basically the sequence analysis methods to predict miRNA-mRNA pairs algorithm is based on sequence complementarity between the mature miRNA and the target site. TargetScan and miRanda are two popular methods of predicting miRNA-mRNA pairs. TargetScan is a computational method applied to predict miRNA target sites conserved among orthologous 3’ untranslated regions (UTRs) of vertebrates [50]. This method requires a 6-nt or 7-nt match to the seed region of the miRNA (nucleotide 2-8). The miRanda algorithm, which also searches for complementarity matches between miRNAs and 3’ UTRs using dynamic programming alignment, considers that the interaction is probably not simple hybridization by optimal base pairing [51]. So miRanda algorithm not only do sequence-matching to assess first whether two sequences are complementary and possibly bind, but also calculate free energy to estimate the energetics of this physical interaction and use evolutionary conservation as an informational filter (determine if an miRNA matches an mRNA in more than one species). We will take the union of the miRNA lists predicted by TargetScan or miRanda for a specific mRNA and use the lists as all the potential miRNAs that could target this mRNA. Then we do CKE modeling to narrow the predictions by checking the curve fitting.

5.2 Previous Microarray Data Analysis

In publication [2], the microarray data of miRNAs were interpolated from 4 points into 7 points, and were centered the values about their mean, and performed principal components analysis and k-means clustering. The classification results give 3 dominant patterns: Go-Up, Transient, and Go-Down: Class 1, exhibiting a downward trend; Class 2, transient in nature; and Class 3, exhibiting an upward trend. To evaluate the effect of GCNF -/-
on these time patterns, [2] examined the F-statistic for the interaction of GCNF-/- on the time effects. 200 probes were identified where the GCNF treatment had a significant effect on the time pattern of expression. From figure 5.2 [2], we can see that 13 miRNAs of class 1 are predicted by TargetScan to target 5 key mRNAs regulating the ES differentiation, in which case the expression levels of these miRNA-mRNAs pairs are positively correlated. 7 miRNAs of class 3 are predicted by TargetScan to target the same 5 key mRNAs, in which case the expression levels of these miRNA-mRNA pairs are negatively correlated. From figure 5.2 [2], 13 miRNAs of class 1 are predicted by TargetScan to target 5 key mRNAs regulating the ES differentiation, in which case the expression levels of these miRNA-mRNAs pairs are positively correlated. 9 miRNAs of class 3 are predicted by TargetScan to target the same 5 key mRNAs, in which case the expression levels of these miRNA-mRNA pairs are negatively correlated. We found that among the 31 miRNA-mRNA pairs predicted by TargetScan and 38 pairs by miRanda, only 7 pairs are in common.

5.3 Previous Biological Interpretation of correlation analysis

To characterize how miRNAs are linked to the regulatory networks of ES cells self-renewal and differentiation, publication [2] classifies miRNAs into 3 classes.

Class 1 miRNAs have high expression level on days 0-1 and low expression level on day 6.

Class 3 miRNAs have low expression level on days 0-1, and high expression level on day 6.

Class 2 miRNAs gathers all other types of transient evolution from day 0 to day 6.

In our microarray recordings for ES cell differentiation, we have 105 miRNAs of class 1, 78 miRNAs of class 3 and 46 miRNAs of class 2.

Recall that the key genes regulating ES cells include \{GCNF, Oct4, Nanog, Sox2, Esrrb, cMyc, Klf4, Tbx3, Ezh1, Ezh2, Eed\}. These 11 key regulatory genes involve (see [2]) first
Figure 5.1: Dramatic time ordered patterns of differential expression are revealed by miRNA microarray data after retinoic-acid (RA) treatment [2]. 3 distinct classes of miRNAs were identified in [2] in the ES time course.
Figure 5.2: miRNA-mRNA pairs predicted by TargetScan.
Figure 5.3: miRNA-mRNA pairs predicted by miRanda.
the orphan nuclear receptor GCNF (germ cell nuclear factor), or the other interchangeable name NR6A1 (nuclear receptor 6A1), which is the best characterized transcriptional repressor of Oct4 and Nanog. Both Oct4 protein and Nanog protein are the transcriptional factors for two groups of mRNAs: the self renewal regulators (Sox2, Klf4, Esrrb, Tbx3 cMyc), and the differential inhibitors (Ezh1, Ezh2, Eed).

There are 26 miRNAs of class 1 and 23 miRNAs of class 3 which are predicted by reference tables TargetScan or miRanda to target the key genes involved in regulating ES cells. The authors of [2] discussed on the positive or negative correlation between miRNAs and mRNAs, and no specific conclusion was reached in [2] about the potential targets and regulations in ES cell differentiation for the 46 miRNAs of Class 2.

The conclusions of [2] sketch potential regulatory loops for the WT context 5.4 and KO context 5.5. GCNF protein represses the expression of Oct4 and Nanog in WT context, while in KO context the expression of Oct4 and Nanog are weakly repressed because the GCNF protein is knocked out. Oct4 and Nanog protein are activators for class 1 miRNAs, self-renewal regulators and differentiation inhibitors. In the same time Oct4 and Nanog are repressors for class 3 miRNAs. These interactions between (Oct4, Nanog) and other genes are weakened in the WT context because (Oct4, Nanog) expression levels are lower in WT than in KO. Both class 1 and 3 miRNAs repress the self-renewal regulators and differentiation inhibitors, but class 1 miRNAs also repress (Oct4, Nanog) and hox cluster, while class 3 miRNAs repress GCNF. The interactions between class 1 miRNAs and other genes are weakened in WT context because expression levels of class 1 miRNAs are lower in WT than KO (consistent with expression of Oct4 and Nanog), while interactions between class 3 miRNAs and other genes are weakened in KO context because expression levels of class 3 miRNAs are lower in KO than in WT (negatively correlated with Oct4 and Nanog).
Figure 5.4: **Regulatory loops for WT stem cells**. The arrow indicates activation while bar with a hash at end indicates repression. The inhibition sign indicates that activation or repression is weakened or blocked.

### 5.4 Open Questions Left Unresolved in Previous Study

From figure 5.4 and 5.5, hypotheses were proposed that some class 1 miRNAs probably repress (Oct4, Nanog), self-renewal regulators, differentiation inhibitors and Hox cluster.
However, it is natural to ask whether all class 1 miRNAs target a specific gene, like Oct4 as indicated in the figure 5.4 or only some of class 1 target Oct4. We have similar question about class 3 miRNAs, too. Although we can get a predicted list of miRNAs for a specific mRNA G by TargetScan or miRanda, it is still not clear that whether a predicted miRNA
would directly cleave the mRNA $G$ or repress the translation of $G$. We want to know which miRNA in class 1 plays the role of degrading of the target mRNA and which miRNA plays the role of inhibiting the translation of its target mRNA.

Linear correlation analysis does not give any positively correlated relation or negative relation of class 2 miRNAs with other increasingly expressed or decreasingly expressed genes, which implies that no direct possible activation or repression guesses could be made. So we also want to know if class 2 miRNAs possibly have important functions in the ES cells differentiation regulation.

The following are the summarized list of questions we are interested in:

- For a fixed mRNA $G$, find out what specific miRNAs probably directly cleave it.
- For a fixed protein $P$, find out what specific miRNAs probably repress the translation.
- Do class 1 or 3 play a more important role than class 2 do in ES cell differentiation or class 2 miRNAs also involve in ES cell differentiation as well.
Chapter 6

Basic Architectural Motifs for Regulatory Loops

6.1 the Main Interaction Motifs to Model the Impact of MiRNAs on the Regulatory Loops of Differentiation

The databases miRanda and TargetScan(5.0) predict for each miRNA "M" a list TARG(M) of the mRNAs targeted by M: we are interested in the key regulatory genes of ES cells which belong to TARG(M); for instance, "mmu-miR-186" potentially targets Oct4. To validate the reality and impact of such potential interactions, we will apply below our modeling techniques to validate more precisely which miRNAs may possibly repress a given key regulatory gene G, and to determine whether such miRNAs directly degrade their mRNA target G and/or repress the G protein.

Note that the correlation techniques and PCA analysis used in [2] could only provide fairly qualitative indications about such questions. Figure 5.4 gives a global network involving...
interactions of key genes and miRNAs in ES differentiation. However, modeling such a
global network by ODEs generates a large number of parameters, in which case it is not
possible to obtain reliable estimations with this few data points available (19 data points
in WT and 19 data points in KO for each gene). And artificial models, instead of CKE
models derived from chemical reaction laws, may have to be applied in order to include all
reactants for modeling in a large regulation network. We intend to study more precisely
about the different functions of miRNAs on mRNA and proteins, so we consider two basic
architectures of small motifs in the following sections.

6.2 Motif A Architectures Linking miRMNAs and the ES
      Cells Regulatory Network

We will first study a family of potential interaction motifs, each one of which involves the
interactions between one miRNA "M" with one key regulatory gene G targeted by M, and
with proteins which are potential transcriptional activators of G.

More generally, denoting by G any fixed "downstream" mRNA, we want to find the most
likely upstream miRNAs repressing the expression of G, taking into account of potential
transcriptional and post-transcriptional factors. These small groups of interacting factors
define analogous interacting architectures (or "motifs"), which we will generically denote
by Motif A architectures.
6.2.0.1 A Typical Example of Motif A

Figure 5.4 and 5.5 indicate that GCNF is a transcriptional repressor of the mRNA Oct4, and that class 1 miRNAs may target and degrade Oct4. And according to [5, 8], it is plausible to select Oct4 protein and Nanog protein as transcriptional activators of the mRNA Oct4. And mmu-mir-186 is an miRNA predicted by miRanda to target Oct4. Thus we have an example of motif A (figure 6.2).

6.3 Key Families of Motif A Architectures

Among all potential miRNAs indicated by miRanda or TargetScan to target the mRNA Oct4, we wanted to identify those which directly downregulate the expression of Oct4. Below, we model the family of potential Motif A architectures involving arbitrary potential
downregulation pairs (M,G) where "M" is any one of our 266 miRNAs and "G" is any one of the 10 key regulatory mRNAs displayed in figure 1, namely Oct4, Nanog, Sox2, Klf4, Esrrb, Tbx3, cMyc, Ezh1, Ezh2, Eed. We naturally impose that mRNA \( G \) must belong to the list of mRNAs targeted by \( M \), according to either the miRanda or TargetScan data bases (We take the union of the miRNAs list predicted by miRanda or TargetScan). 19 predicted miRNAs for Oct4, 2 predicted miRNAs for Nanog, 29 predicted miRNAs for Sox2, There are 238 pairs of (M,G) in all.

Based on [2][12], (Oct4, Nanog) both have a transcriptional repressor GCNF, and potential transcriptional activators (Oct4, Nanog, Sox2) [5]. We will want to determine which combinations of these 3 activators is the most probable (given our data), in the downregulation of a fixed mRNA \( G \), for instance Oct4, by a given miRNA \( M \). So we will separately
model the impact, in the previous down regulation of Oct4 by $M$ of each one of the following seven transcriptional activators combinations: (Oct4), (Nanog), (Sox2), (Oct4, Nanog), (Oct4, Sox2), (Nanog, Sox2), (Oct4, Nanog, Sox2). So there could be seven possible motif A architectures involving Oct4 and M. Since there are 19 miRNAs predicted to target Oct4 and 2 miRNAs predicted to target Nanog, then there are $7(19+2)=147$ motifs for these 2 downstream mRNAs.

For (Sox2, Klf4, Esrrb, Tbx3, cMyc, Ezh1, Ezh2, Eed), [2] suggests (see figure 5.4) that (Oct4, Nanog) are transcriptional activators or repressors. This analysis identifies 217 pairs of (M,G) for these 8 downstream mRNAs. Thus in all there are $217+147=364$ motifs of type A that need to be validated by model fitting.

### 6.4 Motif B Architectures Linking miRMNAs and the ES Cells Regulatory Network

For each fixed downstream protein $P$, let $G$ be the associated mRNA producing the protein $P$. The upstream miRNAs inhibit the translation of $G$ and repress the expression of $P$. This defines a motif type denoted as Motif B.

![Figure 6.3: Motif B](image)

We have generated a set of Western blot data for 4 proteins, namely GCNF, Oct4, Nanog, Sox2. Hence in this paper we will restrict the study of Motifs B architectures to
6.4 Figure 6.4: Oct4 protein is the downstream factor, the upstream factors are the mRNA Oct4 that is the producer of the protein and the miRNA/miRNAs that inhibits the translation.

Situations where protein $P$ is one of these 4 proteins. Figure 8 shows an example for motif B, we fix the downstream protein Oct4, and select 3 miRNAs (mmu-mir-290, mmu-mir-296, mmu-mir-138) from the predicted list by TargetScan and miRanda as the repressers. We want to find out what miRNAs have the major influence on these 4 proteins from the predictions from miRanda or TargetScan. We will try at most 3 miRNAs as upstream factors for each of proteins Oct4, Nanog and Sox2, and only 1 miRNA as upstream factor for protein GCNF, for GCNF is knocked-out in KO context and the number of data points are too few to estimate parameters of the models with 2 or 3 upstream factors. This defines 5337 architectures of motif B.

6.5 Modeling and Validation Methodology for Motif A and Motif B architectures

We have introduced 2 basic regulatory motifs to describe the molecular interactions of miRNAs with their targeted mRNAs and the associated proteins. To quantify the validity of potential motif A or motif B architectures, we will model these architectures by parametrized chemical kinetics equations and evaluate the quality of fit of these models to
Our motifs modeling will select the motif A and motif B architectures having high levels of fit with the microarray data. To reach robust conclusions we will apply a "parameter parsimony" principle, and among the validated motif architectures, we will favor those having the smallest number of parameters. The combinatorial possibilities then unavoidably impose the study of a large number of such motifs, as explained below.

Our explicit parameterization of chemical modeling with associated quality of fit computations yields quantitative results validating the prediction of the miRNAs which are most likely to downregulate any specific mRNA or to inhibit the translation of any given mRNA. Our approach involves algorithmic parameterization of nonlinear chemical kinetics models, and goes further than the well established linear analysis by PCA and correlation techniques. This parsimonious modeling methodology combined with detailed quality of fit evaluations provides a useful complement to classical linear data mining techniques and to the targeting information provided by well known sequence-matching methods.
Chapter 7

Modelling Basic Interaction Motifs by Chemical Kinetics Equations

7.1 Modeling Microarray Data by Chemical Kinetics Equations

In the present study, we will use several types of chemical kinetics equations (CKEs), which are parametrized nonlinear ordinary differential equations (ODEs) to model the mechanism of interactions between miRNAs, their target mRNAs and the proteins associated to these targets. The interactions mainly involve cis-regulation, transcription, post-transcription, and translation. J. Goutsias and his collaborators [3,9] have proposed and applied in other contexts several efficient types of nonlinear chemical kinetics equations to model transcription and cis-regulation.

In this work, we modify the CKEs introduced in [3,9] for transcription and cis-regulation, in order to take into account the repressive impact of miRNAs, and we also derive a chemical
kinetics equation for translation, involving the influence of repressive miRNAs during post-transcription.

Chemical kinetics equations (CKEs) are widely used to model many biological processes or biochemical reaction networks for predicting, simulating or analyzing the expression levels dynamics of chemical species. Many fundamental processes in cellular biology, such as regulator binding, transcription, translation and degradation can be modeled by CKEs at the molecular level.

In published literature on this topic, CKEs have been formulated to describe the pathway structure for mRNAs and proteins [9,35,79,80], which have the following form:

$$\frac{dg(t)}{dt} = -\beta g(t) + \kappa F(\theta, p_1(t), \ldots, p_k(t))$$  \hspace{1cm} (7.1)

where $g(t)$ and $[p_1(t), \ldots, p_k(t)]$ represents the concentrations of a specific mRNA gene $G$, and proteins $[P_1, \ldots, P_k]$ which are transcription factors of the gene $G$. Here the parameters $\beta > 0$ and $\kappa > 0$ are the degradation rate and transcription rate of gene $G$, and $\theta$ is a vector of parameters such as reaction rates or other constants coming from reduction of the modeling derivation. The nonlinear function $F(\theta, p_1(t), \ldots, p_k(t))$ models how the transcription factors $[P_1, \ldots, P_k]$ repress or activate the mRNA gene $G$. Several types of regulation function $F(t)$ have been proposed and applied [35,36], namely

1. customized functions
2. sigmoid functions
3. step functions
The often used Hill function $Hill(p, \theta, n)$ as a function of concentration $p$, parameters $\theta$ and $n$ is as follows:

$$Hill(p, \theta, n) = \frac{p^n}{\theta^n + p^n}$$

It is based on sigmoid functions, but customized functions can also be derived or justified by applying adequate chemical kinetics principles, such as the law of mass action or the Michaelis-Menten law. The variables involved in the CKEs can be concentrations or relative expression levels of molecules such as proteins, mRNAs, miRNAs and so forth. The parameters of the CKEs, such as degradation rates, transcription rates, translation rates, equilibrium constants and so forth, are usually unknown and are often difficult to measure experimentally with enough accuracy.

Mathematical modeling by nonlinear chemical kinetics equations generates complicated parameter estimation problems from recorded microarray data.

### 7.2 Chemical Kinetics Equation for Motif A

Select arbitrarily one mRNA gene "G" and one miRNA gene "M". Call "P" the protein generated by the gene "G". There are essentially two main modalities of interaction within the triplet of molecules ["G", "P", "M"], and we now model these (potential) interactions by chemical kinetics equations linking the expression levels of these 3 molecules.

The regulatory motif A is a small size interaction model describing how the rate of change for the expression of "G" (a downstream mRNA) depends on the expression levels of its upstream factors, which include the post-transcriptional repressor miRNA "M", and two sets of proteins rep(G) and act(G), namely the transcriptional repressors and transcriptional activators of "G".
7.2.1 Post-transcriptional Repressors and Architectural Interaction Motif A

The regulatory architectural "motif A" is a small size interaction model describing how the rate of change for the expression of "G" (a downstream mRNA) depends on the expression levels of its upstream factors, which include the post-transcriptional repressor miRNA "M", and two sets of proteins rep(G) and act(G), namely the transcriptional repressors and transcriptional activators of "G".

To model the transcription process involving interactions between these transcription factors (proteins) and their downstream mRNA "G". We introduce a nonlinear chemical kinetics equation (CKE) similar to the equation proposed in [3,9], but with a complementary term encoding the repressive influence of miRNA "M" on its target mRNA "G".

Denote by \( g(t) \), \( p(t) \) and \( m(t) \) the expression levels of molecules "G", "P" and "M" at time \( t \). We call \( \{ R_1, R_2, \ldots \} \) and \( \{ A_1, A_2, \ldots \} \) the proteins belonging respectively to rep(G) and act(G). Denote by \( r_i(t) \) and \( a_j(t) \) the respective expression levels of proteins \( R_i \) and \( A_j \).

We thus model motif A by the following chemical kinetics equation:

\[
\frac{dg(t)}{dt} = -\beta g(t) - vg(t)m(t) + \kappa F(t)
\]  

(7.2)

where \( \beta > 0 \) is the degradation rate of \( G \), \( v > 0 \) is the reaction rate between \( G \) and \( M \), \( \kappa > 0 \) is the transcription rate, and \( F(t) \) is the fraction of DNA templates committed to transcription of the mRNA gene \( G \). The fraction \( F(t) \) is modeled by setting

\[
RF_i(t) = \frac{1}{(1 + u_ir_i(t))^S_R_i} \quad AF_j(t) = \frac{1}{(1 + w_ja_j(t))^S_A_j}
\]

(7.3)

\[
REP(t) = \prod_{R_i \in \text{rep}(G)} RF_i(t) \quad ACT(t) = 1 - \prod_{A_j \in \text{act}(G)} AF_j(t)
\]

(7.4)

\[
F(t) = REP(t)ACT(t)
\]

(7.5)
Here $SR_i > 0, u_i > 0$ and $SA_j > 0, w_j > 0$ are respectively the number of binding sites and the affinity constant for the transcriptional factors $R_i$ and $A_j$.

Note that the transcription repressors $R_i$ combine multiplicatively their individual impacts $RF_i(t)$ in $F(t)$; a similar remark applies to the impacts of the transcription activators.

The term $\kappa F(t)dt$ is the concentration of new $G$ molecules synthesized by transcription during the small time interval $[t, t + dt]$.

In the same time interval, the repressive interactions of molecules $M$ and $G$ eliminates $vg(t)m(t)dt$ molecules of $G$, and natural decay destructs $\beta g(t)dt$ molecules of $G$.

The specific form (7.3) of the fraction $F(t)$ of DNA templates committed to the transcription of gene $G$ has been studied in other contexts by [3, 9]. We give below, in the last paragraph of this chapter, the main arguments and assumptions which justify the expression of $F(t)$ in our context.

### 7.3 Chemical Kinetics Equation for Translation Repressors

#### 7.3.1 Interaction Motif B

The regulatory architectural "motif B" is a small size interaction model describing how the rate of change for the expression of a downstream protein $P$ depends on the expression level of its upstream factors, which include the mRNA gene $G$ producing protein $P$ and the set $\text{rep}(G) = [M_1, M_2, \ldots]$ of miRNA repressing the translation of $G$. The respective concentrations at time $t$ of protein $P$, miRNA genes $M_i$, and mRNA gene $G$ are denoted by $p(t), m_i(t), g(t)$. 
7.3.2 CKEs to model motif B architectures

We model motif B interaction architectures by the following chemical kinetics equation, which modify the CKEs in [3, 7] by complementary terms:

\[ \frac{dp}{dt} = -\gamma p(t) + \lambda g(t)H(t) \]  \hspace{1cm} (7.6)

\[ RP_i(t) = \frac{1}{(1 + u_i m_i(t))^{SM_i}} \]  \hspace{1cm} (7.7)

\[ H(t) = \prod_{M_{i \text{rep}(G)}} RP_i(t) \]  \hspace{1cm} (7.8)

where \( \gamma > 0 \) and \( \lambda > 0 \) are resp. the degradation rate and the translation rate for protein \( P \).

The parameters \( SM_i > 0 \) and \( u_i > 0 \) are resp. the number of binding sites and the affinity constant driving the repressive impact of miRNA \( M_i \) on the mRNA gene \( G \). The global repressive impact of all the miRNA molecules \( M_i \) on the translation of \( G \) is encoded in \( H(t) \), which decreases when the miRNA concentrations \( m_i(t) \) increases.

The term \( H(t) \) is the fraction of \( G \) molecules committed to the translation.

7.4 Biochemical Assumptions and CKEs Derivation for Motifs A and B

In the main CKE of motif A (see (7.2)), the complementary term \( -v g(t)m(t) \) encodes the down-regulation of the target mRNA "G" by miRNA "M", under the following 2 fairly natural biochemical assumptions

- Assumption 1: Although there maybe many miRNAs that can bind to the target "G", this
architectural motif tentatively assumes that "M" is the only miRNA which can strongly bind at some specific site of gene "G", and that this binding then causes the degradation of the corresponding molecule "G".

- Assumption 2: Once a molecule of "G" and a molecule "M" actually bind at this specific site, then this "G"-molecule degrades in a very short time interval, before binding within any other "M" molecule.

Under these assumptions, we have the following reaction:

\[ M + G \rightarrow O \quad (7.9) \]

where the molecules "O" are the degradation outputs and have concentration \( o(t) \). The Law of Mass Action in chemical kinetics states that the rate at which a chemical is generated is proportional to the product of the concentrations of the reactants with a proportionality constant \( v > 0 \), so that \( \frac{do(t)}{dt} = vg(t)m(t) \). This explains the corresponding term \(-vg(t)m(t)\) in the CKE (7.2) of motif A providing the expression \( \frac{do(t)}{dt} \).

To derive the form of the functions \( F(t) \) and \( H(t) \) in the respective CKEs of motif A and motif B, we will introduce more assumptions beyond the Assumptions 1 and 2.

The main concepts and rigorous derivations needed to support the form we have adopted for the function \( F(t) \) in the CKE for motif A are quite similar to the essential ideas required to derive the proper form of \( H(t) \) in the CKE of motif B.

Hence we will now focus only on the rigorous derivation of the translation CKE for motif B, which introduces the main ideas and concepts.

Recall first that in [3], the translation process of a gene "G" generating the protein "P" is driven by the following very basic chemical kinetics equation

\[ \frac{dp(t)}{dt} = -\gamma p(t) - \lambda g(t) \quad (7.10) \]
where $\gamma$ and $\lambda$ are respectively the degradation rate and the translation rate of protein $P$.

Here, we need to take into account the complementary repressive influence of the set $\text{rep}(G) = [M_1, M_2, \ldots]$ of miRNA inhibiting the translation of $G$. We call $m_j(t)$ the concentration of $M_j$-molecules at time $t$. We make the following fairly natural assumptions applied for motif B:

Assumption 3: If a specific molecule of gene "G" binds with any one of the miRNA repressors $M_j$ listed in $\text{rep}(G)$, then this G-molecule will fail to translate.

Assumption 4: As long as a molecule of gene "G" has bound with none of the miRNA repressors listed in $\text{rep}(G)$, then the translation of this specific G-molecule can initiate freely.

Assumption 5: Each miRNA repressor $M_j$ of gene $G$ can only bind with gene $G$ at $S_j$ specific sites; call $BIND_j$ this set of $S_j$ specific binding sites. We assume that the sets $BIND_1, BIND_2, \ldots, BIND_k$ of binding sites are pairwise disjoint.

Assumption 6: For any given "G"-molecule, call $X_j$ the random number of sites in $BIND_j$ which actually bind with "$M_j$"-molecules. We assume that the $k$ random variables $X_1, \ldots, X_k$ are independent random variables.

The term $\lambda g(t)$ in (7.10) is the concentration of protein "P"-molecules synthesized per unit of time when all the molecules of gene $G$ are committed to the translation of "G". However, at any time $t$, only a fraction of the existing "G"-molecules $t$ are committed to the translation of "G", due to the inhibiting action of the miRNA repressors of gene $G$.

So we need compute the fraction of "G"-molecules which are actually committed to the translation into protein "P"-molecules.

Suppose for the moment that $k = 1$, i.e. that there is only one miRNA gene $M = M_1$ in the list $\text{rep}(G)$ of miRNAs repressing the mRNA gene $G$. Call $S = S_1 = \text{cardinal}(BIND_1)$
the total number of possible binding sites for the miRNA gene $M = M_1$ on the binding region of "G"-molecules.

For $0 \leq s \leq S$, denote by $G_s$ the set of "G"-molecules which have exactly $s$ binding sites actually bound to "M"-molecules. At time $t$, call $g_s(t)$ the concentration of "$G_s$"-molecules, and $m(t)$ the concentration of miRNA "M".

We have the following forward and backward reactions:

$$G[s] + M \rightarrow G[s + 1]$$ \hfill (7.11)

$$G[s + 1] \rightarrow G[s] + M$$ \hfill (7.12)

Call $m_b(t)$ the concentration of free "M"-molecules which bind on the "G[s]"-molecules to produce "G[s+1]"-molecules in the forward reaction 7.11. By molecular collision theory, $m_b$ is proportional to the product of the total concentration $m = m_1$ of "M"-molecules by the concentration of vacant binding sites $(S - s) \cdot g_s$, so that, for some constant $c_f$, we have

$$m_b(t) = c_f(S - s)m(t)g_s(t)$$ \hfill (7.13)

Call $m_f(t)$ the concentration of miRNAs freed by the backward reaction 7.12. Again by molecular collision theory, $m_f$ is proportional to the concentration $(s + 1)g_{s+1}$ of "M"-molecules actually bound to one of the $(s + 1)$ occupied binding sites of the "G"-molecules belonging to $G[s + 1]$, so that for some constant $c_b$ we have

$$m_f(t) = c_b(s + 1)g_{s+1}(t)$$ \hfill (7.14)

At chemical equilibrium, we have naturally $m_f = m_b$, and hence

$$c_f(S - s)m(t)g_s(t) = c_b(s + 1)g_{s+1}(t)$$
Letting $u = c_f/c_b$, we obtain
\[
g_{s+1}(t) = u[(S - s)/(s + 1)]m(t)g_s(t)
\]
By an easy recurrence on $s$, this implies
\[
g_s(t) = C_S^s[um(t)]^sg_0(t) \tag{7.15}
\]
where the $C_S^s = \frac{(S!)}{s!(S-s)!}$ are the classical binomial coefficients. So the total concentration $g(t)$ of "G"-molecules can be expressed by
\[
g(t) = \sum_{s=S}^{\infty} g_s(t) = g_0(t) \sum_{s=0}^{S} C_S^s[um(t)]^s = g_0(t)[1 + um(t)]^S \tag{7.16}
\]
By our assumptions, if at least one of the $S = S_1$ binding sites of a "G"-molecule is actually bound to an "M"-molecule, then this "G"-molecule will fail to translate, so the fraction $H(t)$ of "G"-molecules committed to the translation into protein "P"-molecules is clearly given by $H(t) = \frac{g_0(t)}{g(t)}$.

In view of the expression (7.16) just obtained we thus have in the case $k = 1$ of a single miRNA repressor for gene G.
\[
H(t) = \frac{1}{[1 + um(t)]^{S_1}} = \frac{1}{[1 + um_1(t)]^{S_1}} \tag{7.17}
\]
So the simple CKE (7.10) for translation must be replaced in our context, and when $k = 1$, by
\[
\frac{dp}{dt} = -\gamma p(t) + \lambda g(t)H(t) = -\gamma p(t) + \lambda g(t)\frac{1}{[1 + um_1(t)]^{S_1}} \tag{7.18}
\]
where $p(t)$ is the concentration of protein $P$, $g(t)$ is the concentration of mRNA gene $G$ and $m_1(t)$ is the concentration of miRNA $M = M_1$.

By recurrence on the number $k$ of miRNA repressors in the list $rep(G) = \{M_1, M_2, ..., M_k\}$, and using the Assumptions 5 and 6 described above, we can extend the argument just
given for the case \( k = 1 \), and prove that in the generic case where \( k \geq 1 \), then the fraction

\[
H(t) = \frac{g_0(t)}{g(t)}
\]

of "G"-molecules committed to the translation into protein "P"-molecules is given by:

\[
H(t) = \prod_{i=1}^{k} \frac{1}{(1 + u_i m_i(t))^{s_i}} \tag{7.19}
\]

As we pointed out above, quite similar ideas enable us to derive the form of the function \( F(t) \) and the CKE of motif A.

### 7.5 Simplified Chemical Kinetics Equations for High Degradation Rate

If there is no protein data available, consider the following equation for translation process with the assumption that the miRNA repression for the protein levels is ignorable [3]:

\[
\frac{dp(t)}{dt} = -\delta p(t) + \lambda g(t) \tag{7.20}
\]

In equation (7.20), \( p(t) \) is the concentration of a specific protein \( n \) at time \( t \), \( g(t) \) is the concentration of mRNA \( G \), \( \lambda \) is the translation rate of the mRNA \( G \), and \( \delta \) is the degradation rate of protein \( P \).

We can see that this equation for the rate of change of protein is different from the equation we have introduced above. This is a simplified chemical equation proposed by Goutsias and ignore the influence by the miRNAs.

Equation (7.20) is easy to solve:

\[
p(t) = \exp(-\delta t)p(0) + \int_{s=0}^{s=t} \lambda \exp(-\delta(t-s))g(s) \tag{7.21}
\]
We interpolated the data into 19 time points (time unit = 8 hours), the above can be approximated as:

\[ p(t) = \exp(-\delta t)p(0) + \frac{1}{2} \cdot \lambda \sum_{s=0}^{t-1} [\exp(-\delta(t-s))g(s) + \exp(-\delta(t-s-1))g(s+1)] \]  (7.22)

Then

\[
p(t) = \exp(-\delta t)p(0) + \frac{1}{2} \cdot \sum_{s=0}^{t-1} [\exp(-\delta(t-s))g(s) + \exp(-\delta(t-s-1))g(s+1)]
\]

\[
\leq \exp(-\delta t)p(0) + 1/2 \lambda g(t) + \sum_{s=0}^{t-1} [\exp(-\delta(t-s))g(s)]
\]

\[
= \exp(-\delta t)p(0) + 1/2 \lambda g(t) + 
\lambda [\exp(-\delta)g(t-1) + \exp(-\delta \cdot 2)g(t-2) + ... + \exp(-\delta t)g(0)]
\]

\[
= \exp(-\delta t)p(0) + \lambda(0.5g(t) + \exp(-\delta)g(t-1) + ... + \exp(-\delta t)g(0))
\]

If \( \exp(-\delta)g(t-1) < 5\% \cdot 0.5g(t) \), then \( p(t) \sim \exp(-\delta t)p(0) + 0.5 \cdot \lambda g(t) \). For the 11 key genes we studied in this paper, \( \max(g(t-1)/g(t)) < 1.7 \) when \( t > 0 \). So

\[
\exp(-\delta) \cdot g(t)/g(t-1) < \exp(-\delta) \cdot 1.7 < 0.05 \cdot 0.5
\]

which gives \( \delta > 4.22 \), which means the half-life of the protein should be less than \( \ln(2)/4.22 \cdot 8 = 1.32 \) hours (note that the time unit in our equations are 8 hours). Therefore, if \( \delta > 4.22 \), 
\[ p(t) \sim \exp(-\delta t)p(0) + \lambda 0.5g(t) \]. When

\[
\lambda g(t) >> \exp(-deltat)p(0)
\]  (7.23)

is satisfied, for \( t > 0 \), we have

\[ p(t) \approx \lambda \cdot 0.5g(t), t > 1 \]  (7.24)

The above equation means \( p(t) \) is approximately proportional to \( r(t) \), i.e. \( p(t) = \alpha g(t) \), where \( \alpha \) is a constant. This result holds under simplified ODE modeling the translation.
process, when the associated proteins of the key 11 genes have protein half-life within 1.32 hours and (7.23) is satisfied.

### 7.6 Formal Invariance of our CKE models by Scale Changes

In most microarray data sets, the "absolute" expression levels of molecules are not chemically meaningful since they are measured via optical analysis of fluorescence intensities, but the main biologically meaningful quantities are the relative expression levels between arbitrary pairs of recorded chemical species. This is one of the reasons why the classical graphic displays of microarray data by "heat maps" actually involve logarithms of the data.

Since we want to apply our chemical kinetics equations to microarray data sets for which the most meaningful values are ratios of expression levels data, we will look at how the chemical kinetics equation changes if we apply generic linear transformations to the expression data.

Denote by $g(t), p(t), m(t)$ the respective chemical concentrations of mRNA gene $G$, its associated protein $P$, and miRNA gene $M$ targeting $G$. Assume that $g(t), p(t), m(t)$ are linked with the concentrations $r_i(t)$ and $a_j(t)$ of resp. repressive and activating factors $R_i(t) \in rep(G), A_j(t) \in act(G)$, by the following "motif A" CKE

$$\frac{dg(t)}{dt} = -\beta g(t) - vg(t)m(t) + \kappa REP(t)ACT(t)$$

(7.25)
where

\[ REP(t) = \prod_i RF_i(t) \]  \hspace{1cm} (7.26)

\[ ACT(t) = 1 - \prod_j AF_j(t) \]  \hspace{1cm} (7.27)

\[ RF_i(t) = \frac{1}{(1 + u_ir_i(t))^{SR_i}} \]  \hspace{1cm} (7.28)

\[ AF_j(t) = \frac{1}{(1 + w_ja_j(t))^{SA_j}} \]  \hspace{1cm} (7.29)

Assume now that the actually recorded expression levels of these molecules, namely \( \hat{g}, \hat{p}, \hat{m}, \hat{r}_i, \hat{a}_j \) are linked to the true concentrations \( g, p, m, r_i, a_j \) by unknown but constant scale changes, \( \gamma, \mu, \rho_i, \alpha_j \) so that we have for all times \( t \) the linear relations

\[ \hat{g}(t) = g(t)/\gamma, \quad \hat{p}(t) = p(t)/\tau, \quad \hat{m}(t) = m(t)/\mu \]  \hspace{1cm} (7.30)

\[ \hat{r}_i(t) = r_i(t)/\rho_i, \quad \hat{a}_j(t) = a_j(t)/\alpha_j \]  \hspace{1cm} (7.31)

Define new parameters by the explicit formulas

\[ \hat{\beta} = \beta \]  \hspace{1cm} (7.32)

\[ \hat{v} = v\mu \]  \hspace{1cm} (7.33)

\[ \hat{\kappa} = \kappa/\gamma \]  \hspace{1cm} (7.34)

\[ \hat{u}_i = \rho_i u_i \]  \hspace{1cm} (7.35)

\[ \hat{w}_j = \alpha_j w_j \]  \hspace{1cm} (7.36)

Then transferring these last two sets of linear relations into the CKE linking the true concentrations, we obtain the following new CKE where only recorded expression levels are involved,

\[ \frac{d\hat{g}(t)}{dt} = -\hat{\beta}\hat{g}(t) - \hat{v}\hat{g}(t)\hat{m}(t) + \hat{\kappa}REP(t)ACT(t) \]  \hspace{1cm} (7.37)
where

\[ \widehat{REP}(t) = \prod_i \widehat{RF}_i(t) \quad \widehat{ACT}(t) = 1 - \prod_j \widehat{AF}_j(t) \] (7.38)

\[ \widehat{RF}_i(t) = \frac{1}{(1 + \hat{u}_i\hat{r}_i(t))^{SR_i}} = RF_i(t) \] (7.39)

\[ \widehat{AF}_j(t) = \frac{1}{(1 + \hat{w}_j\hat{a}_j(t))^{SA_j}} = AF_j(t) \] (7.40)

Note that the algebraic form of this new CKE linking the actual recorded expression levels of is exactly the same as the algebraic form of the CKE linking the true concentrations, but of course with new unknown parameters. The new unknown parameters are related to the original unknown parameters by scale changes but this has no practical impact, since true concentrations are not accessible through microarray data.

Assume now that each actually recorded gene or protein expression level \( y(t) \) is linked by some linear relation of the type \( x(t) \rightarrow y(t) = ax(t) \) to the true concentration \( x(t) \) of the same gene or protein, where the scale change coefficient can freely depend on the specific gene or protein considered, but is constant in time.

Our preceding argument shows then that each biochemical hypothesis positing a CKE model of motif A type between true concentrations is strictly equivalent to assuming that a similar CKE model of motif A type also holds between recorded expression levels.

Similarly computations easily show that the same global invariance of the models by multiple scale changes is true for CKE models of motif B type. Actually for CKE models of type B, the global invariance of these CKE models also hold for more general affine relations of the form. \( x(t) \rightarrow y(t) = ax(t) + b \) between expression levels \( y(t) \) and true concentrations \( x(t) \).

One can also prove that the same global invariance by general affine transformations will hold for a slight algebraic extension of the CKE models of motif A type. After such multiple affine transformations are applied to true concentrations, the new unknown parameters...
are explicitly linked to the unknown original parameters by non linear relations which are explicit rational fractions. Of course these relations between parameters have no practical impact.

The key theoretical result is that we can freely normalize linearly our recorded expression levels, using different normalizing constants for each recorded gene, before estimating the parameters of our CKE models of motif A types or of motif B types. Since our Western Blot protein data are also essentially linearly linked to the corresponding unknown absolute concentrations, the preceding arguments also applies to cover the case of the actual protein data we have digitized from Western Blots experimental acquisitions.

The preceding global invariance result will also enable us to select an adequate distance between dynamic profiles of recorded expression levels.
Chapter 8

Parameter Estimation for the CKE models of Motifs A and B

8.1 Parameter Estimation for Nonlinear CKEs

8.1.1 Fitting Chemical Kinetics Dynamic Systems to Data

For actual nonlinear CKE modeling of biochemical processes, estimating the parameters involved in these CKEs is a major challenge. Model size strongly increases the complexity of parameter estimation, and can involve very large amounts of computing time in our context, due to the huge number of potential genes interaction architectures. In our context, massive modeling by nonlinear CKE and estimation of their unknown parameters is not feasible "blindly" due to the magnitude of the number of CKEs (more than 30,000 kinetics equations) and the combinatorial complexity for the selection of directly interacting pairs of genes. Moreover the CKEs we introduce to model the interactions
between miRNAs and mRNAs are strongly nonlinear and involve 5 to 10 unknown parameters per CKE. Parameter estimation in such systems is of course strongly influenced by measurement errors in microarray data.

8.1.2 Model Sensitivity to Errors in Experimental Data

Errors in expression levels measurements are inevitable in microarray data, due to the complexity of cellular environments. These errors in measurements are systematically estimated and listed by most technical providers of microarray data.

A natural question is then to evaluate the associated uncertainty on the estimated parameter values computed from experimental data, as well as model sensitivity to changes in parameters. To avoid model instability, regularization techniques are used to control the impact of the noise corrupting the recorded data [13, 42]. Bayesian methods [41], which can make good use of estimated standard deviations of recorded expression levels, provide approximate probability distributions for parameter values. But Bayesian methods such as Markov Chain Monte Carlo (MCMC) techniques, are still too difficult to implement and/or computationally too expensive when the problem is high dimensional [43].

8.2 Parameter Estimation by Cost Functions Minimization

A predominant strategy for parameter estimation in systems of CKEs or ODEs is to minimize a cost function evaluating the discrepancy between model predictions and experimental data.

The concrete goal is to find the parameter values which optimize the quality of fit between
simulated time-course for concentrations (or expression levels) and associated recorded observations.

In published applications, optimization methods usually start from an initial guess which involves more or less intensive global searches in the parameter space. To get a good initial guess, generic stochastic minimization methods such as artificial genetic algorithms, simulated annealing, Markov Chain Monte Carlo (MCMC) have been used as global search techniques [38, 39], as well as ”pattern search” methods [8, 9] or linear code parameter search [6].

The genetic algorithm (GA) is a search heuristic that mimics the process of natural evolution. This heuristic is routinely used to generate useful solutions to optimization and search problems. The basic concept of GAs is designed to simulate processes in natural system necessary for evolution, specifically those that follow the principles first laid down by Charles Darwin of survival of the fittest.

Simulated annealing is a generalization of a Monte Carlo method for examining the equations of state and frozen states of n-body systems [38]. It works by emulating the physical process whereby a solid is slowly cooled so that when eventually its structure is ”frozen”, this happens at a minimum energy configuration.

Markov Chain Monte Carlo (MCMC) method is a class of algorithms for sampling from probability distributions based on constructing a Markov chain that has the desired distribution as its equilibrium distribution. MCMC uses the previous sample values to randomly generate the next sample value, generating a Markov chain as the transition probabilities between sample values are only a function of the most recent sample value.

Pattern search (PS) refers to a family of numerical optimization methods that do not require the gradient of the problem to be optimized and PS can hence be used on functions that are not continuous or differentiable. They varied one theoretical parameter at a time.
by steps of the same magnitude, and when no such increase or decrease in any one parameter further improved the fit to the experimental data, they halved the step size and repeated the process until the steps were deemed sufficiently small \[59\]. A linear code is a subspace of a finite dimensional vector space over a finite field. The linear code parameter search algorithm (LCPS) was developed out of the need to survey high dimensional parameter spaces in which varying one or two parameters at a time, while keeping the others fixed, is simply not feasible. It has at its core a (not necessarily linear nor binary) code with efficient covering properties of the high dimensional parameter space \[6\]. More specifically, it is essential to find out the minimum covering radius of a code. Combined with such global search results, local methods such as gradient descent techniques have also been used to minimize these types of cost functions, but mostly for smaller scale problems involving much smaller sets of CKEs.

### 8.2.1 An example of parameter estimation techniques for systems of CKEs

For instance, in paper \[10\], CKEs with 9 equations were formulated for a chemical system of reactions. These 9 equations involve 113 parameters. 31 experiments were performed with variations in the initial concentrations of three molecules and the time intervals, which provides \(9 \times 31\) equations.

So the ratio of the number of data points to the number of parameters is \(279/113 \approx 2.47\) which is quite low, and this is certainly an undesirable feature from a purely statistical point of view.

Then the parameter estimation problem was converted into an optimization problem with
loss function

\[ \text{Loss}(\text{PAR}) = \sum_{j=1}^{31} w_j ||z_j(t) - J(\text{PAR}, t)_j||^2 \]

Where \( w_j = 1/||z_j||^2 \) is the scalar weight, \( z_j(t) \) is the measured data for the \( j^{th} \) experiment at time \( t \), and the norm is weighted square norm in \( L^2 \) space.

The minimization problem was solved by a combined use of global and local optimization techniques. Global searches of the admissible space of parameters were performed by the MATLAB routines ”ga” and ”patternsearch” which implement artificial genetic algorithms and pattern search algorithms to provide suitable candidates as initialization values for subsequent local searches.

In [10], these local searches were implemented by the MATLAB code ”fmincon”, which encodes a sequential quadratic programming algorithm. As the name implies, sequential quadratic programming (SQP) methods are iterative methods solving at each iteration a quadratic programming problem (QP), which is a linearly constrained optimization problem with a quadratic objective function [60].

The approach in [10] still required the use of published bounds defining the adequate ranges for the unknown parameters.

8.2.2 Parameter Estimation Strategy Adopted in Our Study

Stochastic minimization algorithms typically require large amounts of computing time, a drawback which becomes prohibitive when the number of unknown parameters is high, and gradient descent methods strongly depend on their initialization points. So generic methods for cost minimization are generally less efficient than specifically designed strategies adapted to a specific form of cost function.
We have hence developed our own algorithms to estimate the parameters of our CKEs based models from microarray data and to then compute the quality of fit between models and data. In our work, this quality of fit, adequately balanced by strongly favoring parsimoniously parametrized models, becomes an essential clue to decide which potential interactions we should validate between miRNAs, mRNAs, and the associated proteins. Reducing the combinatorial complexity by focusing on biologically meaningful architectural motifs is another essential step we have adopted to increase the ratio of ”number of data points” over the ”number of unknown parameters”. Thus we have focused on a key genes known to be involved in ES cells differentiation, such as GCNF, Oct4, Nanog, Sox2, Klf4, Esrrb, cMyc, Tbx3, Ezh1, Ezh2, Eed, and on pairs of miRNA and mRNA genes which are listed as potentially interacting in the published tables Targetscan and miRanda.

8.3 Parsimony in Parameters Identification

Our CKEs based models are build and parametrized to enable a robust discrimination between actual interacting pairs of between miRNA and mRNA genes, and non interacting pairs of miRNA and mRNA genes.

To increase the robustness of the model, and according to major statistical results due to Vapnik [S2], one should increase the ratio of the the number of data points over the number of unknown parameters in the model.

Consider the following equation:

$$\frac{dg(t)}{dt} = \kappa F(t) - v_m(t)g(t) - \beta g(t)$$
Where the fraction function $F(t)$ is:

$$F(t) = \prod_{i \in \text{rep}(G)} \frac{1}{(1 + u_ir_i(t))^S_{R_i}} \prod_{j \in \text{act}(G)} \frac{1}{(1 + w_ja_j(t))^S_{A_j}}$$

Denote $n_A$ to be the number of activators, $n_R$ be the number of repressors, then the number $PAR$ of unknown parameters we need to estimate for each such CKE is $PAR = 3 + 2(n_A + n_R)$. Let $T$ be the number of data points available for each one of the recorded expression levels $g(t)$, $m(t)$, $a_i(t)$ and $r_j(t)$, $t = t_1, t_2, \ldots, t_N$. Then for each mRNA, these $T$ data generate $T$ constraints on the unknown parameters of the corresponding CKE.

The Vapnik results show us that good accuracy of parameter estimates require fairly high values of the ratio of $T/PAR$. So $PAR \ll T$ is a necessary constraint. If $PAR > T$, then the models are generally overfitted and the parameters are poorly estimated.

Since there are only 38 data points after interpolation for WT and KO, this yields the following strong constraint on the number of parameters, namely $n_A + n_R \leq 3$.

### 8.4 Estimation of Parameters by Gradient Descent

Here we take a simple model for example. For a fixed protein $P$ and upstream miRNA $M$, the equation to solve is the following:

$$\frac{dp(t)}{dt} = \lambda g(t) \frac{1}{(1 + um(t))^S} + \gamma p(t) \quad (8.1)$$

Suppose miRNA $M$ is a repressor candidate of protein $P$, and suppose that $\gamma$, $u$ are fixed, then we want to minimize the following function:

$$Z(\lambda, \gamma, u, S) = \sum_t [p(t) - p(0) + \int_0^t (\lambda p(t) - \gamma g(t) \frac{1}{(1 + um(t))^S})^2 \quad (8.2)$$

where $\int_0^t$ is a notation for mid-point Riemann sum on interval $[0, t]$. We tested the gradient descent method to solve the minimization problem. First of all, we want to pick up a "good"
initial value of parameters \((\gamma, \lambda, u, S)\), usually by crude search or using the Matlab functions such as "ga". It would be better to find out the range of the parameters before applying global search. However degradation rate \(\gamma\) and \(\lambda\) is unaccessible for most proteins. And since there are only a few fraction of all mRNAs that have more than one binding sites for miRNAs and only \(0.02\%\) mRNAs have as many as 7 binding sites \([63]\), so it is reasonable for us to restrict \(0 \leq S \leq 7\).

If \(\frac{1}{(1+um(t))} \leq 0.05\), then it means that only 5% of all the mRNAs are committed to translate the proteins, i.e., over 95% of the mRNAs are bound by the miRNAs. From the findings in \([63]\) that the miRNA does not act like a genetic switch, and that the miRNAs play a general role of fine-tuning in gene expression, so we do not expect that there are too many mRNAs bound by the miRNAs, which would cause an dominated effect to gene expression. Thus we can restrict \(u\) by the following:

\[
\max \left( \frac{1}{(1+um(t))^S} \right) > 0.05
\]

and solve the upper bound of \(u\), denoted as \(u_{max}\).

With \(\gamma > 0\), \(\lambda > 0\), \(u \in [0, u_{max}]\), \(S \in [1, 7]\), we applied the Matlab function "ga" or "patternsearch" and get an initial vector of parameters \([\gamma_0, \lambda_0, u_0, S_0]\). To implement gradient descent, we set up the cost function \(Z(\gamma, \lambda, u, S)\):

\[
Z(\gamma, \lambda, u, S) = \sum_t (p(t) - p(0) + \int_0^t \gamma p(t) - \lambda^{(k)} g(t) \frac{1}{(1+um(t))^S})^2
\]  

(8.3)
Then we take the partial derivative of the function $Z(\gamma, \lambda, u, S)$.

\[
\frac{dZ}{d\gamma} = 2 \sum_t (p(t) - p(0) + \int_0^t \left( \lambda p(t) - \gamma g(t) \frac{1}{(1 + um(t))^S} \right) g(t) \frac{1}{(1 + um(t))^S})
\]

\[
\frac{dZ}{d\lambda} = 2 \sum_t (p(t) - p(0) + \int_0^t \left( \lambda p(t) - \gamma g(t) \frac{1}{(1 + um(t))^S} \right) \left[ (\lambda S m(t) \frac{1}{(1 + um(t))^{S+1}}) \right])
\]

\[
\frac{dZ}{du} = 2 \sum_t (p(t) - p(0) + \int_0^t \left( \lambda p(t) - \gamma g(t) \frac{1}{(1 + um(t))^S} \right) (\lambda S m(t) \frac{1}{(1 + um(t))^{S+1}})) (- \int_0^t \lambda \log\left( \frac{1}{(1 + um(t))^{S+1}} \right) \frac{1}{(1 + um(t))^S})
\]

\[
\frac{dZ}{dS} = 2 \sum_t (p(t) - p(0) + \int_0^t \left( \lambda p(t) - \gamma g(t) \frac{1}{(1 + um(t))^S} \right) (\lambda S m(t) \frac{1}{(1 + um(t))^{S+1}})) (- \int_0^t \lambda \log\left( \frac{1}{(1 + um(t))^{S+1}} \right) \frac{1}{(1 + um(t))^S})
\]

Let $\nabla Z = (\frac{dZ}{d\beta}, \frac{dZ}{d\kappa}, \frac{dZ}{du}, \frac{dZ}{dS})$, For the $n^{th}$ iteration of gradient descent:

1. take $h_3 = 1$, 

   \[
   (\beta^{(n+1)}, \kappa^{(n+1)}, u^{(n+1)}, S^{(n+1)}) = (\beta^{(n)}, \kappa^{(n)}, u^{(n)}, S^{(n)}) - h_3 \nabla Z^{(n)}
   \]

2. While $Z^{(n+1)} \geq Z^{(n)}$, let $h_3 = h_3/2$, then renew the value of of $Z^{(n+1)}$ until $Z^{(n+1)} < Z^{(n)}$, if $h_3 < TOL$, where $TOL$ is a very small number, stop the iteration and keep $(\beta^{(n)}, \kappa^{(n)}, u^{(n)}, S^{(n)})$ as the solution.

3. If any parameter turns out to be negative, set it to be zero; If $S^{(n+1)} > 10$, take $S_{(n+1)} = 10$.

4. Renew the value of $Z^{(n+1)}$, if $Z^{(n+1)} < Z^{(n)}$, continue, if not, stop with the $(\beta^{(n)}, \kappa^{(n)}, u^{(n)}, S^{(n)})$ as the solution.

### 8.5 Parameter Estimation Algorithm for the CKEs of Motifs A and B

There are 2 types of equations we use to model the 2 basic motifs:

\[
\frac{dp(t)}{dt} = -\beta p(t) + \kappa g(t) H(t)
\]
where

\[ H(t) = \prod_{M_i \in \text{rep}(G)} \frac{1}{(1 + um_i(t))^{S_i}} \]

and

\[ \frac{dg(t)}{dt} = -\beta g(t) - vg(t)m(t) + \kappa F(t) \]

where

\[ F(t) = \prod_{i \in \text{rep}(G)} \frac{1}{(1 + u_i r_i(t))^{S_{R_i}}} \left( 1 - \prod_{j \in \text{act}(G)} \frac{1}{(1 + w_j a_j(t))^{S_{A_j}}} \right) \]

where \( g(t), m(t), r_i(t), a_j(t) \) and \( p(t) \) are known for \( t = 1, \ldots, 19 \).

For simplicity and not losing generality, we consider the integrated (by discrete approximation) equation of motif A with only one upstream factor:

\[ g(t) - g(0) = -\beta \int_0^t g(s)ds - v \int_0^t m(s)g(s)ds + \kappa \int_0^t F(s)ds \]

where \( t = 0, 1, \ldots, 19 \) and

\[ F(t) = \begin{cases} \frac{1}{(1 + up(t))^S}, & \text{case of repression} \\ 1 - \frac{1}{(1 + up(t))^S}, & \text{case of activation} \end{cases} \]

The idea of our method is simple: if we already know the value of affinity constant \( u \) and binding sites \( S \), we can estimate the values of decay rate \( \beta \) and transcription rate \( \kappa \) by solving a simple constrained linear programming problem. Note that the factor \( q = 1/(1 + up(t)) \) is in the range of interval \([0, 1]\). We can do an exhaustive search on \( q \), set the precision to be 0.01, and let \( u = \text{average}_t((1 - q)/(qp(t))) \). And \( S \) is an integer in the range of \([1, 10]\). Given each value of \( u \) and \( S \), we can compute the value of \( \int_0^t F(s)ds \), thus, there are linear equations:

\[ g(t) - g(0) = -\beta \int_0^t g(s)ds - v \int_0^t g(s)m(s)ds + \kappa \int_0^t F(s)ds \]
Let
\[ \epsilon = \max_t |(t) - g(0) + \beta \int_0^t g(s)ds + v \int_0^t g(s)m(s)ds - \kappa \int_0^t F(s) | \]

To minimize \( \epsilon \), we can solve the following constrained linear programming problem:

\[
\begin{align*}
\text{minimize} \quad & \epsilon \\
\text{subject to} \quad & -\epsilon \cdot 1 < C \cdot x < \epsilon \cdot 1
\end{align*}
\]

where \( 1 \) is a \( n \times 1 \) vector, \( n = 19 \) in this case, \( x = [\beta, v, \kappa] \), and

\[ C = \left( \begin{array}{ccc} a & b & c \end{array} \right) \]

where \( a = [-\int_0^1 g(s), -\int_0^2 g(s), \ldots, -\int_0^t g(s)]^T \), \( b = [-\int_0^1 g(s)m(s), \ldots, -\int_0^t g(s)m(s)]^T \), \( c = [-\int_0^1 F(s), \ldots, -\int_0^t F(s)]^T \).

Then we will select the best solution of the parameters \( \{u, S, \beta, v, \kappa\} \) to minimize \( \epsilon \).

Consider a more general case for a fixed mRNA \( G \) as the downstream factor, which is activated by protein \( P_1 \), repressed by protein \( P_2 \), and degraded by miRNA \( M \), then we have the following equation:

\[ \frac{dg}{dt} = -\beta g(t) - vg(t)m(t) + \kappa F(p_1(t), p_2(t)) \]

where \( g(t) \) is the expression level of downstream mRNA \( G \), \( p_1(t) \) is the concentration of \( P_1 \), \( p_2(t) \) is the concentration of \( P_2 \), \( m(t) \) is the concentration of \( M \). And

\[ F(p_1(t), p_2(t)) = (1 - \frac{1}{(1 + u_1 p_1(t))^{S_1}})(1 - \frac{1}{(1 + u_2 p_2(t))^{S_2}}) \]

In this example, we need to search the values of \( u_1, S_1, u_2 \) and \( S_2 \), then solve the parameters \( \beta, v \) and \( \kappa \) by constrained linear programming.

Generally, for these two type of equations, we first do the exhaustive search for the affinity constants \( u_i \) and and number of binding sites \( S_i \) for all \( i \in rep(G) \cup act(G) \), and then apply
the constrained linear programming to estimate the other parameters. This algorithm is relatively fast in computing, on laptop PC it usually takes less than 10 minute to estimate a 9-parameter model, depending on the precision tolerance the user choose for searching. The global exhaustive search is for small number of parameters: integer-valued binding sites $S_i$, which take less than 10 values, and at most 3 affinity constants $u_i$ in a small interval $[0,1]$ and can even take bigger searching steps, such as 0.02 or 0.05 depending on the number of parameters. It combines a global search method in reasonable precision and constrained linear programming method, which is easily implemented with high-quality results. Another advantage of this algorithm is that range of the parameter values are not required to provide good estimation results, which is very useful for many of these molecular reaction rates are unaccessible in literature.

8.6 Quality of Fit between Model Predictions and Expression Levels Data

For a specific motif A or motif B, after algorithmic parametrization of the model in order to achieve the best fit to the two sets of microarray data, we have to evaluate how well the parametrized model predicts the observed dynamics for the concentration of the downstream factor.

$$D(t) = \text{measured expression level of the downstream factor at time } t$$

After actual parametrization of the model, let

$$\hat{D}(t) = \text{model-predicted expression of the downstream factor at time } t$$
To assess the quality of fit between model prediction of $\hat{D}(t)$ and recorded microarray data $D(t)$, it would seem natural to compute the relative error of prediction at time $t$ as follows:

$$ERR(t) = \frac{|D(t) - \hat{D}(t)|}{D(t)}$$

However, the relative error of prediction $ERR(t)$ becomes meaninglessly large whenever $D(t)$ is close to zero. To avoid these spurious large values, particularly when $\hat{D}(t)$ visually agrees quite well with $D(t)$ for all $t$, we define the smoothed relative error of prediction to be:

$$SERR(t) = \begin{cases} 
ERR(t) & \text{if } D(t) > 0.15 \cdot \text{mean}(D) \\
\frac{|D(t) - \hat{D}(t)|}{\text{mean}(D)} & \text{if } D(t) < 0.15 \cdot \text{mean}(D)
\end{cases}$$

where mean($D$) is the average of $D(t)$ over all values of $t$. Then we define the global error of prediction $err$ of the model by

$$err = \max_t (SERR(t))$$

The error of prediction $err$ quantifies the accuracy level of the parametrized model, and is percentage valued.

We will say that the model is "acceptable" or "validated" if, for both the WT and the KO microarray data, the "model error" $err$ is less than 10%, or if $err$ is inferior to the relative standard deviation of the recorded expression levels.

### 8.7 Sensitivity of Model Predictions to Small Changes in Parameters Values

Prediction of expression levels by model of motif type A or B could be possibly very sensitive to the values of changes for parameters. We here choose only $\epsilon = 5\%$ as the error
tolerances of the parameters, and see numerically how the model predictions vary when we vary one parameter and fix the others. Denote the parameter we select to change to be $\text{PAR}_0$, and we randomly choose 50 values by uniform distribution, denoted as $\text{PAR}_i$ for $i = 1, \ldots, 50$, from interval $\text{PAR}_0 \cdot [1 - \epsilon, 1 + \epsilon]$ and simulate 50 curves for each value $\text{PAR}_i$. Denote the original model error computed from $\text{PAR}_0$ as $\text{err}_0$, and the maximum model error out of the 50 simulations as $\text{err}_{max}$. The results show that the CKEs of motif type A or B are not much sensitive to parameters. We take a look at an example of motif A with parameters $(\beta, \kappa, v, u_1, u_2, u_3, S_1, S_2, S_3)$ as follows: downstream mRNA=Oct4 miRNA = mmu-mir-186
transcription repressor= GCNF
transcription activators= Oct4, Nanog

This example has a very good model prediction with model error $err_0 = 0.05$ and $0.08$ respectively for WT and KO. We choose $5\%$ as the error tolerance, and select parameter $\beta$. Then we get $err_{max} = 0.13$ for WT simulation, and $err_{max} = 0.15$ for KO simulation (see figure 8.2). Figure 8.3 shows that when $\kappa$ changes in the error tolerance of $5\%$, $err_{max} = 0.08$ and $0.17$ respectively for WT and KO simulations. Figure 8.4 shows that when $v$ changes in the error tolerance of $5\%$, $err_{max} = 0.18$ and $0.14$ respectively for WT and KO simulations. Figure 8.5, 8.6 and 8.7 show that when $u_1or u_2or u_3$ changes in the error tolerance of $5\%$, the values of $err_{max}$ are almost stable respectively for WT and KO simulations.

For $(S_1, S_2, S_3)$, they all equal to $1$, which they are number of binding sites, are integers. So first we shift up by $1$ for each of these three parameters and keep the other two fixed, and see from Figure 8.8, 8.9, 8.10 that $err_{max}$ for these three cases are no bigger than $0.18$, in which the model fitting is not bad.

Parameter estimation gives us $S_1 = S_2 = S_3 = 1$ in this example. When we shift down by $1$ for each of these three parameters and keep the other two fixed, then we have three cases:

- $S_1 = 0, S_2 = S_3 = 1$, which means protein GCNF is excluded as the repressor for mRNA Oct4, then $err_{max} = 2.34$ for WT. Figure 8.11 shows this prediction fits the measured data very badly.

- $S_2 = 0, S_1 = S_3 = 1$, which means protein Oct4 is excluded as the activator for mRNA Oct4, then $err_{max} = 0.13$ for KO. Figure 8.12 shows this prediction fits the measured data fairly well.

82
• $S_3 = 0, S_2 = S_1 = 1$, which means protein Nanog is excluded as the activator for mRNA Oct4, then $\text{err}_{\text{max}} = 1.32$ for KO. Figure 8.13 shows this prediction fits the measured data very badly.

![Figure 8.2: Display of predictions for mRNA Oct4 of motif A when parameter $\beta$ changes for WT and KO.](image)

The downstream factor has upstream proteins GCNF, Oct4, Nanog, and miRNA mmu-mir-186. Blue line = measured expression level, red dash line=predicted expression level by originally estimated parameters, green dotted lines=predicted expression levels by modifying the $\text{PAR}_0$ to be $\text{PAR}_i$. 

83
Figure 8.3: Display of predictions for mRNA Oct4 of motif A when parameter $\kappa$ changes for WT and KO. The downstream factor has upstream proteins GCNF, Oct4, Nanog, and miRNA mmu-mir-186. Blue line = measured expression level, red dash line=predicted expression level by originally estimated parameters, green dotted lines=predicted expression levels by modifying the $PAR_0$ to be $PAR_i$.

8.8 Sensitivity of Parameter Estimates to Errors on Expression Levels

In microarray data, recorded expression levels are corrupted by background "noise" , and the microarray data sets actually report estimated values of the standard deviation for the error of measurement on each recorded expression level. We have undertaken the evaluation of the corresponding uncertainty on estimated parameter values computed from experimental data. This is a highly intensive computing task since each one of the more
Figure 8.4: Display of predictions for mRNA Oct4 of motif A when parameter \( v \) changes for WT and KO. The downstream factor has upstream proteins GCNF, Oct4, Nanog, and miRNA mmu-mir-186. Blue line = measured expression level, red dash line=predicted expression level by originally estimated parameters, green dotted lines=predicted expression levels by modifying the \( PAR_0 \) to be \( PAR_i \).

than 10,000 instances of motif A or motif B architectures we had retained for algorithmic parametrization potentially requires a separate computer analysis to quantify the impact of measurement errors on the estimated parameters. This massive computerized validation is not yet completed, so here we simply present some of the main steps involved in its implementation.

A first approach we have tested is to simulate random perturbations of the recorded expression levels for a specific downstream or upstream factor, where the simulations adds white noise errors to each relevant recorded expression level value, with the adequate standard deviation actually attached to each measurement reported in the microarray data set. We
Figure 8.5: Display of predictions for mRNA Oct4 of motif A when parameter \( u_1 \) changes for WT and KO. The downstream factor has upstream proteins GCNF, Oct4, Nanog, and miRNA mmu-mir-186. Blue line = measured expression level, red dash line=predicted expression level by originally estimated parameters, green dotted lines=predicted expression levels by modifying the \( PAR_0 \) to be \( PAR_i \).

then re-estimate the CKE model parameters, but using these artificially perturbed data . The re-estimation steps are the following.

1. Select a downstream (or upstream) factor FA in a specific CKE model of motif A or motif B type. Simulate 50 randomly perturbed evolution profiles of the factor FA, as we did in section 3.4, and keep the expression levels of all other downstream or upstream factors unchanged.

2. Keep fixed all the values of affinity constants \( u_i, w_j \) and number of binding sites \( SR_i, SA_j \) associated to the various repressors and activators involved in the CKE
model. These fixed values are pre-estimated from the original recorded expression levels, as explained above.

3. Then for each one of the 50 simulated perturbed profile of FA, we re-estimate all the other parameters by our fast constraint linear programming.

4. Call $PAR_0$ the parameters of the CKE model estimated from the original recorded microarray data. Let $PAR_k$ be the parameters (of this CKE model) generated by re-estimation from the $k^{th}$ simulated perturbed dataset.

5. Compute the empirical mean values $MEANPAR$ and standard deviations $STDPAR$.
Figure 8.7: **Display of predictions for mRNA Oct4 of motif A when parameter $u_3$ changes for WT and KO.** The downstream factor has upstream proteins GCNF, Oct4, Nanog, and miRNA mmu-mir-186. Blue line = measured expression level, red dash line = predicted expression level by originally estimated parameters, green dotted lines = predicted expression levels by modifying the $\text{PAR}_0$ to be $\text{PAR}_i$.

of the random re-estimated vector $\text{PAR} = [\text{PAR}_1, \ldots, \text{PAR}_5, 0]$.

Note that in our approach for parameter estimation, we perform first an exhaustive search for the affinity constants and number of binding sites, and then implement a constraint linear programming algorithm to estimate the other parameters. Fixing the values of affinity constants and of the number of binding sites strongly reduces the computing time of each parameter estimation, but provides nevertheless a good exploration of the robustness to noise for our constraint linear programming algorithm.
Figure 8.8: Display of predictions for mRNA Oct4 of motif A when parameter $S_1$ changes for WT and KO. The downstream factor has upstream proteins GCNF, Oct4, Nanog, and miRNA mmu-mir-186. Blue line = measured expression level, red dash line = predicted expression level by originally estimated parameters, green dotted lines = predicted expression levels by modifying the $PAR_0$ to be $PAR_i$.

8.8.0.0.1 Example 1: Robustness Study for a motif A model. We consider the following motif A.

$G =$ downstream factor = mRNA gene Oct4

$M =$ miRNA degrader = mmu-mir-186

$R_1 =$ transcription repressor = GCNF

$A_1 =$ transcription activator = protein Oct4

$A_2 =$ transcription activator = protein Nanog

This CKE model involves 9 parameters ($\beta, \kappa, v, u_1, w_1, w_2, SR_1, SA_1, SA_2$). All affinity constants ($u_1, w_1, w_2$) and number of binding sites ($SR_1, SA_1, SA_2$) are deliberately kept
Figure 8.9: Display of predictions for mRNA Oct4 of motif A when parameter $S_2$ changes for WT and KO. The downstream factor has upstream proteins GCNF, Oct4, Nanog, and miRNA mmu-mir-186. Blue line = measured expression level, red dash line = predicted expression level by originally estimated parameters, green dotted lines = predicted expression levels by modifying the $PAR_0$ to be $PAR_i$.

We choose $G$ as the factor $FA$ to be submitted to 50 random perturbations at each time point, and we re-estimate 50 vectors of parameter values $PAR_i, i = 1, \ldots, 50$ for the vector $[\beta, \kappa, v]$.

The initial estimates computed from the original recorded expression levels downstream are

$$\beta_0 = 0.091, \quad \kappa_0 = 414, \quad v_0 = 0.001$$
Figure 8.10: **Display of predictions for mRNA Oct4 of motif A when parameter $S_3$ changes for WT and KO.** The downstream factor has upstream proteins GCNF, Oct4, Nanog, and miRNA mmu-mir-186. Blue line = measured expression level, red dash line = predicted expression level by originally estimated parameters, green dotted lines = predicted expression levels by modifying the $PAR_0$ to be $PAR_i$.

After 50 simulated perturbations of the $G$ expression levels and corresponding re-estimations for these 3 parameters, we obtained 50 estimated values for each parameter, namely $\beta_i$, $\kappa_i$ and $v_i$, for $i = 1, \ldots, 50$. Then we computed the empirical means and standard deviations for each one of these 3 parameters:

\[
\text{MEAN}_\beta = 0.097, \quad \text{STD}_\beta = 0.091
\]

\[
\text{MEAN}_\kappa = 439, \quad \text{STD}_\kappa = 117
\]

\[
\text{MEAN}_v = 0.001 \quad \text{STD}(v) = 6 \times 10^{-4}
\]
Figure 8.11: **Display of predictions for mRNA Oct4 of motif A when parameter $S_1$ changes for WT and KO.** The downstream factor has upstream proteins GCNF, Oct4, Nanog, and miRNA mmu-mir-186. Blue line = measured expression level, red dash line = predicted expression level by originally estimated parameters, green dotted lines = predicted expression levels by modifying the $PAR_0$ to be $PAR_i$.

8.8.0.0.2 **Example 2: Robustness study for a motif A model.** We consider the following motif A.

$G =$ downstream factor $=$ mRNA gene Sox2

$M =$ miRNA degrader $=$ mmu-mir-21

$A_1 =$ transcription activator $=$ protein Oct4, $A_2 =$ transcription activator $=$ protein Nanog

This model involves 7 parameters ($\beta, \kappa, v, w_1, w_2, SA_1, SA_2$).
Figure 8.12: Display of predictions for mRNA Oct4 of motif A when parameter $S_2$ changes for WT and KO. The downstream factor has upstream proteins GCNF, Oct4, Nanog, and miRNA mmu-mir-186. Blue line = measured expression level, red dash line = predicted expression level by originally estimated parameters, green dotted lines = predicted expression levels by modifying the $P AR_0$ to be $P AR_1$.

All affinity constants ($w_1, w_2$) and number of binding sites ($SA_1, SA_2$) are fixed. We choose $G$ as the factor $FA$ to be perturbated and we implemented 50 random perturbations of the recorded expression levels of $G$. We then generated 50 re-estimated vectors of parameters values $PAR_i$, $i = 1, \ldots, 50$ for $\beta, \kappa$ and $v$. The initial estimation computed from the original recorded expression levels downstream are

$$\beta_0 = 0.08, \quad \kappa_0 = 380.00, \quad v_0 = 6 \times 10^{-6}$$
Figure 8.13: Display of predictions for mRNA Oct4 of motif A when parameter $S_3$ changes for WT and KO. The downstream factor has upstream proteins GCNF, Oct4, Nanog, and miRNA mmu-mir-186. Blue line = measured expression level, red dash line=predicted expression level by originally estimated parameters, green dotted lines=predicted expression levels by modifying the $PAR_0$ to be $PAR_i$.

The empirical means and standard deviations computed from 50 re-estimated values of these 3 parameters were the following.

$\text{MEAN}_\beta = 0.05$, $\text{STD}_\beta = 0.04$

$\text{MEAN}_\kappa = 263$, $\text{STD}_\kappa = 158$

$\text{MEAN}_v = 7 \times 10^{-6}$, $\text{STD}(v) = 1.5 \times 10^{-6}$

8.8.0.0.3 Example 3: Robustness study for a motif B model. We consider the following motif B.
This model involves 8 parameters ($\beta, \kappa, u_1, u_2, u_3, S_1, S_2, S_3$). All affinity constants ($u_1, u_2, u_3$) and number of binding sites ($S_1, S_2, S_3$) are fixed. We simulated 50 randomly perturbed expression levels for the downstream factor $P$, and re-estimated 50 vector values $PAR_i$, $i = 1, \ldots, 50$ for the parameters $\beta, \kappa$. The estimates computed from the original recorded expression levels were the following:

$$\beta_0 = 0.27, \quad \kappa_0 = 9 \times 10^{-5}$$

After 50 random perturbations and simulations and re-estimations for these 2 parameters, we computed the empirical means and standard deviations for the parameter estimates:

$$MEAN_\beta = 0.27, \quad STD_\beta = 0.03$$

$$MEAN_\kappa = 9 \times 10^{-5}, \quad STD_\kappa = 1.3 \times 10^{-5}$$

8.8.0.4 Example 4: Robustness study for a motif B model. We consider the following motif B, quite similar in structure to the preceding one.

$P = \text{downstream factor} = \text{protein Oct4}$

$M_1 = \text{miRNA repressor} = \text{mmu-miR-542-3p}$

$M_2 = \text{miRNA repressor} = \text{mmu-miR-484}$

$M_3 = \text{miRNA repressor} = \text{mmu-miR-138}$

This model involves 8 parameters ($\beta, \kappa, u_1, u_2, u_3, S_1, S_2, S_3$). All affinity constants ($u_1, u_2, u_3$) and number of binding sites ($S_1, S_2, S_3$) are fixed. The initial parameter values (computed
from original data) of $\beta, \kappa$ were

$$\beta_0 = 0.2656, \quad \kappa_0 = 8.6903 \times 10^{-5}$$

After 50 simulated perturbations of the downstream factor $P$, and corresponding re-estimations of the 2 parameters $\beta, \kappa$, we obtained the empirical means and standard deviations of their estimators.

$$MEAN_\beta = 0.27, \quad STD_\beta = 0.03$$

$$MEAN_\kappa = 9 \times 10^{-5}, \quad STD_\kappa = 1.3 \times 10^{-5}$$

8.8.0.0.5 Robustness Evaluations In the above 4 examples, the mean values of the estimated parameters are reasonably close to the initial ones computed from the original data.

The standard deviations are relatively large for the first two examples of motif A, and fairly small for the last two examples of motif B. To explain this, we point out that the standard deviations of measurements errors are of the order of 10% to 20% of the recorded expression levels for the genes G in the two motif A models, while for the protein $P$ data involved in the two examples of model B, we have deliberately selected an optimistic relative value of 5% for the unknown size of measurements errors on protein data.

These examples show that estimated parameters can fluctuate to quite an extent due to the errors of measurements affecting recorded expression levels. For each motif A or motif B architecture we have parameterized, we intend to perform this robustness study to confirm the direct evaluation of the quality of fit. However this second level validation program is quite costly to complete, and is for the moment still a work in progress, due to the very large number of architectures we have parametrized.
Chapter 9

Model Validated Interactions between MicroRNAs and mRNA genes

9.1 Modeling for Basic Motifs of Type A

There are 3 groups of mRNAs we want to study as downstream factors of motif A.

1. (Oct4, Nanog), each of which has transcription repressor GCNF as figure 1. The potential transcription activators of these two mRNAs are proteins (Oct4, Nanog, Sox2) [5].

2. Self-renewal regulators (Sox2, Klf4, cMyc, Tbx3, Esrrb). According to figure 1, we use (Oct4, Nanog) as the transcription activators.

3. Differential inhibitors (Ezh1, Ezh2, Eed). According to figure 1, we use (oct4, Nanog)
as the transcription repressors.

For each downstream factor, the potential upstream miRNA degrader could be selected from the predictions of TargetScan and miRanda. Once we fix the downstream gene and its transcriptional factors and select an miRNA that may target the mRNA from the list of predictions, we can validate whether this hypothesis is plausible by modeling motif A.

Knowing that GCNF is the key repressor for Oct4 and Nanog, for each miRNA predicted from miRanda and TargetScan, we tried the combinations (GCNF, Oct4), (GCNF, Nanog), (GCNF, Sox2), (GCNF, Oct4, Nanog), (GCNF, Oct4, Sox2), (GCNF, Nanog, Sox2), (GCNF, Oct4, Nanog, Sox2) as the upstream TFs for Oct4 and Nanog.

The solution and model quality shows that (GCNF, Oct4, Nanog) is the best combination for both downstream factors Oct4 and Nanog. Together with (GCNF, Oct4, Nanog), mmu-miR-186, 466, 103 and 107 are the miRNAs such that modeling for mRNA Oct4 are plausible. And we present the best model for mRNA Nanog, in which the corresponding miRNA is mmu-miR-139.

We can see in Table 9.1 that most of the validated miRNAs of motif A are in class 2 (transient). The only exception is that all the 12 validated miRNAs for target Ezh2 are in class 3. There is no validated pair for mRNA Nanog or Esrrb, i.e. no miRNA that directly degrades mRNA genes Nanog or Esrrb can be validated by modeling the potential pairs predicted by miRanda or TargetScan. However, our datasets of recorded microarray data contains only 266 known miRNAs, so we did not have the possibility to include in our modeling a small group of miRNAs predicted by miRanda or TargetScan but for which we have no recorded microarray data. Therefore it is possible that more miRNA-mRNA pairs could be validated by motif A models if the available microarray data had included all the identified miRNAs available.
Table 9.3 presents all validated miRNA-mRNA pairs of motif A for the 11 mRNAs. We have evaluated 364 models of motif A and found that 88 out of these 364 models of motif A are validated with model error less than 10%. So there are around 24% motifs of type A that are validated by modeling.

Table 9.1: **Motif A: Number of Validated miRNA-mRNA Pairs**

<table>
<thead>
<tr>
<th>mRNA gene G</th>
<th>miRNAs targeting G</th>
<th>validated in class 1</th>
<th>validated in class 3</th>
<th>validated in class 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oct4</td>
<td>19</td>
<td>6</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Nanog</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Sox 2</td>
<td>29</td>
<td>9</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>Klf 4</td>
<td>44</td>
<td>2</td>
<td>0</td>
<td>0</td>
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<td>Esrrb</td>
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<td>0</td>
<td>0</td>
</tr>
<tr>
<td>cMyc</td>
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<td>12</td>
<td>0</td>
<td>0</td>
</tr>
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<td>18</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
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<td>20</td>
<td>6</td>
<td>0</td>
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**9.2 Modeling of Basic Motif of Type B**

We have determined a list of 5337 models of motif B to be validated for the 4 proteins (Oct4, Nanog, Sox2, GCNF). The computation result shows that there are 51 combinations
Figure 9.1: **Display of a validated motif of type A for Oct4.** The top left and right are profiles of miRNA 186 for WT and KO over day 0-6 respectively. The middle left and right are profiles of transcription factors of Oct4 for WT and KO over day 0-6 respectively. Blue solid line represents the expression evolution of protein Oct4, green dash line represents the expression evolution of protein Nanog, red dotted-solid line represents the expression of protein GCNF. The bottom left and right are profiles of mRNA oct4 for WT and KO respectively. The blue line is the measured expression; the red dash line is the approximated expression predicted by the model with upstream TF GCNF as repressor, (Oct4, Nanog) as activators, and miRNA mmu-miR-186 degrading mRNA Oct4. err represents the model error, std represents the relative standard deviation of the measurement.
Figure 9.2: Display of a motif of type A for nanog. The top left and right are profiles of miRNA 139 for WT and KO over day 0-6 respectively. The middle left and right are profiles of transcription factors of Nanog for WT and KO over day 0-6 respectively. Blue solid line represents the expression evolution of protein Oct4, green dash line represents the expression evolution of protein Nanog, red dotted-solid line represents the expression of protein GCNF. The bottom left and right are profiles of mRNA Nanog for WT and KO respectively. The blue line is the measured expression; the red dash line is the approximated expression predicted by the model with upstream TF GCNF as repressor, (Oct4, Nanog) as activators, and miRNA mmu-miR-139 degrading mRNA Nanog. err represents the model error, std represents the relative standard deviation of the measurement.
Figure 9.3: **Display of a validated motif of type A for Sox2.** The top left and right are profiles of miRNA 21 for WT and KO over day 0-6 respectively. The middle left and right are profiles of transcription factors of Sox2 for WT and KO over day 0-6 respectively. Blue solid line represents the expression evolution of protein Oct4, green dash line represents the expression evolution of protein Nanog. The bottom left and right are profiles of mRNA Sox2 for WT and KO respectively. The blue line is the measured expression; the red dash line is the approximated expression predicted by the model with upstream TF (Oct4, Nanog) as activators, and miRNA mmu-miR-21 degrading mRNA Sox2. err represents the model error, std represents the relative standard deviation of the measurement.
of miRNAs selected from TargetScan or miRanda that are validated for protein Oct4 and 11 miRNAs validated for protein GCNF by modeling motif B. But no plausible models of motif B are found for proteins Nanog and Sox2. Figure 9.4 presents a validated example of protein Oct4 with 3 upstream miRNAs (542-3p, 484, 138) as repressors. Figure 9.5 presents a validated example of protein GCNF with miRNA 181b as repressor.

Table 9.4 shows that the number of validated miRNAs for the 4 proteins. For inhibiting the translation of protein Oct4, there is 1 miRNA validated in class 1, 3 miRNAs validated in class 3, and 5 miRNAs in class 2. For inhibiting the translation of protein GCNF, there is 0 miRNA validated in class 1, 4 miRNAs validated in class 3, and 7 miRNAs in class 2. This result agrees quite well with figure 5.4 and 5.5 which indicates that class 1 miRNAs do not target GCNF but class 3 miRNAs do.

Table 9.5 presents part of the result for motif B (see all list of results for motif B in annex). We list the validated miRNAs for for only Oct4 protein and GCNF protein. Table 4 shows that all the validated miRNAs for protein Oct4 by modeling motif B are the combinations of 3 miRNAs (19 miRNA candidates in all).

Table 9.6 shows that there are 11 validated miRNAs repressing protein GCNF through motif B architectures among 83 miRNA candidates. As we mentioned before, we only have the WT data for protein GCNF, so that the number of data points of protein GCNF is half the data points available for the other proteins. So we select only 1 miRNA as the upstream repressor for protein GCNF because we restrict the number of parameters when we have less data points.
Figure 9.4: **Display of a validated motif of type B for Oct4.** Protein Oct4 and its upstream miRNAs profile for WT and GCNF-KO respectively. The upper left is profile of WT upstream miRNAs. The upper right is profile of KO upstream miRNAs. Blue line is 542-3p expression, green line is 484 expression, red line is 138 expression. In lower graphs, blue line is the measured expression; the red dash line is the approximated expression predicted by the model with upstream the The model error of the WT model and of the KO model are presented below the figures respectively.
Figure 9.5: **Display of a validated motif of type B for GCNF.** Protein GCNF and its upstream miRNAs profile in WT. The upper graph is profile of WT upstream miRNA. Blue line is the measured expression; the red dash line is the approximated expression predicted by the model with upstream the The model error of the WT model is presented below the figures.
9.3 Examples of Invalidated Models of type B for Proteins Nanog and Sox2

Figure 9.6 presents a model of motif B for protein Nanog. We take 2 miRNAs candidates repressing protein Nanog [1], no such combination succeeds to provide a plausible model. This could be due to the lack of enough miRNA candidates in the list that are predicted to target Nanog by miRanda or TargetScan.

For Sox2, there is no validated model of motif B either. All modeling results for protein Sox2 show that the translation rate $\gamma$ is almost zero. So no matter what upstream miRNAs we select, the modeling prediction for Sox2 are the same as figure 9.7. We can see that the degradation rate of Sox2 protein in WT cells looks much more greater than in KO cells. But our modeling made the assumption that the degradation rates in both WT and KO are the same.

Note that the half-life of a protein like Sox2 may quite possibly change in different cell cultures; for instance, protein Oct4 has a half-life of 90 minutes in undifferentiated P19 cells [17] and of 6 to 8 hours in NIH3T3 cells transfected with wild type Oct4 [18].

If we now generate parametrized models of the same motif B architecture, but with the new assumption that the half-life of protein Sox2 is different for WT cells and for KO cells, then we obtain much better model predictions displayed in figure 9.8. The estimated half-life of protein Sox2 is then 20.5 hours for WT cells, and 64.5 hours for KO cells. Although the half-life of protein Sox2 may be very different in WT and KO, which would indeed explain the implausible initial modeling results for Sox2, we still need more explicit biological explanation about the large difference of half-life values.
Figure 9.6: Display of an invalidated motif of type B for Nanog. Protein Nanog and its upstream miRNAs profile for WT and GCNF-KO respectively. The upper left is profile of WT upstream miRNAs mmu-mir-470. The upper right is profile of KO upstream miRNAs mmu-mir-470. In lower graphs, blue line is the measured expression; the red dash line is the approximated expression predicted by the model with upstream the The model error of the WT model and of the KO model are presented below the figures respectively.

9.4 Modeling Hox Cluster

Hox cluster in figure 3 play a very important role in differentiation of ES cells. When Hox cluster are not repressed by (Ezh1, Ezh2, Eed), they activates the ES differentiation; when Hox cluster are repressed by (Ezh1, Ezh2, Eed), then ES differentiation are also repressed.
Figure 9.7: **Display of an invalidated motif of type B for Sox2.** Protein Sox2 and upstream miRNAs mmu-mir-134 and 21. Upper left: WT profiles of upstream miRNAs. Upper right: KO profiles of upstream miRNAs. In lower graphs: blue line = measured expression levels, red dash line = predicted expression levels. Model errors of prediction are too large for validation.

It is natural to think of modeling Hox cluster by chemical kinetics equation of motif type A, in which case (Ezh1, Ezh2, Eed) are transcription repressors. Although the protein expression levels of (Ezh1, Ezh2, Eed) are not available because of technology limitations, it is possible that combinations of the three proteins (Oct4, Nanog, Sox2) are transcription
Figure 9.8: **Display of a validated simplified model for Sox2.** Protein Sox2 profile for WT and GCNF-KO respectively. Blue line is the measured expression; the red dash line is the approximated expression predicted by the model with upstream the The model error of the WT model and of the KO model are presented below the figures respectively.
activators/repressors for the Hox cluster for (Oct4, Nanog, Sox2) regulate a relatively large proportion of genes (Young Lab). It is possible that none of the combinations of the three proteins (Oct4, Nanog, Sox2) are the transcription factors for a fixed mRNA $G_H$ from Hox cluster, which means that both the proteins and miRNA upstream factors are not identified for a downstream factor mRNA $G_H$ of motif A. Since the miRNA is not considered to have big influences on the mRNA expression [1], we can first simplify motif of type A by considering only the interaction of transcription factors and their downstream mRNA.

The following is the chemical kinetics equations involving only interaction of transcription factors and the downstream mRNA [3]:

$$\frac{dg(t)}{dt} = \beta g(t) + \kappa F(t)$$  \hfill (9.1)

Where

$$IF(t) = REP(t)ACT(t)$$

$$RF_i(t) = \frac{1}{(1 + u_iR_i(t))^{SR_i}}, \quad AF_j(t) = \frac{1}{(1 + w_jA_j(t))^{SA_j}}$$

$$REP(t) = \prod_{R_i\in rep(G)} RF_i(t), \quad ACT(t) = 1 - S \prod_{A_j\in act(G)} AF_j(t)$$

For a specific mRNA $G$ in Hox cluster, the upstream transcription factors could be 7 possible combinations of (Oct4, Nanog, Sox2): (Oct4), (Nanog), (Sox2), (Oct4, Nanog), (Oct4, Sox2), (Oct4, Nanog, Sox2). Given 41 mRNAs of Hox cluster, we will have $7 \times 41$ models when transcription factors are activators, $7 \times 41$ models when transcription factors are repressors. We consider the case that two or three transcription factors, a combination we listed above, are either all repressors or all activators for the downstream factor. Because the time course expression levels of these three proteins, by looking at the evolution curves in WT data (Figure 3.16), are strongly positively correlated. Therefore this defines 574
models.

For each model, we do crude search for $\text{mean}[RF_i(t)]$ or $\text{mean}[AF_j(t)]$ on interval $[0,1]$ by a step 0.02 when the number of transcription factors is less than or equal to 2, by a step 0.1 when the number of transcription factors is 3. Then solve the corresponding parameters of $u_i$ or $w_j$. $SR_i$ or $SA_j$ take integer value from 1 to 2 for we assume that there are not many binding sites in order to save computation time. Each model with only one transcription factor takes about 9 seconds for computation time, each model with 2 transcription factors take about 3-4 minutes for computation time, and each model with 3 transcription factors take about 6-10 minutes. So these 574 models involving only interactions between proteins and mRNAs takes around a day to complete computation.

Since we did not take into account of the miRNA influence of degradation, it is natural that we lose the criteria of quality of fit a little bit. Here we say the model is "validated" if the model error $err$ is 20% for these models, or if $err$ is inferior to the relative standard deviation of the recorded expression levels.

The results show that 7 models are validated, they are Hox A9, Hox B7, Hox C10, Hox C13, Hox D3, Hox D8, Hox D9. All these validated model have the same activators (Oct4, Nanog). However we can see from the graphs that, for Hox B7 and Hox D3, the predicted levels by the model are almost straight lines. We check the parameters of Hox B7 and Hox D3 and found that the values of affinity constants $w_1$ and $w_2$ are zeros, i.e. (Oct4, Nanog) do not really effect the expression level for Hox B7 and D3. So we simply exclude these two genes from the validated genes.

The model errors of Hox A9, Hox C13, Hox D8, Hox D9 are all less than 10%, which means the model fitting would be better or at least stay the same if we add one more parameter from the interaction of degradation between mRNA and miRNA. So for these mRNAs we do not have to compute the model fitting of motif type A, because we will always get
validated models.

Then we only have to look at Hox C10, which has 4 predicted miRNAs from the list of targetscan or miRanda. The modeling results show that none of the four miRNAs, which are (mmu-mir-33, mmu-mir-155, mmu-mir-143, mmu-mir-345), can be validated very well with the transcriptional factors (Oct4, Nanog) (See figure 9.16, 9.17, 9.18, 9.19).
Figure 9.9: **Modeling fitting result of Hox B7.** Top left and right are expression levels of Oct4 and Nanog proteins for WT and KO context respectively, where blue line is Oct4 protein, green line is Nanog protein. Bottom left and right are expression levels of mRNA Hox B7 for WT and KO respectively. Blue line is the observed expression level of Hox B7, red dash line is the model prediction of Hox B7. "err"=model error, "std"=standard deviation of measurement.
Figure 9.10: **Modeling fitting result of Hox A9.** Top left and right are expression levels of Oct4 and Nanog proteins for WT and KO context respectively, where blue line is Oct4 protein, green line is Nanog protein. Bottom left and right are expression levels of mRNA Hox A9 for WT and KO respectively. Blue line is the observed expression level of Hox A9, red dash line is the model prediction of Hox A9. "err"=model error, "std"=standard deviation of measurement.
Figure 9.11: **Modeling fitting result of Hox C10.** Top left and right are expression levels of Oct4 and Nanog proteins for WT and KO context respectively, where blue line is Oct4 protein, green line is Nanog protein. Bottom left and right are expression levels of mRNA Hox C10 for WT and KO respectively. Blue line is the observed expression level of Hox C10, red dash line is the model prediction of Hox C10. "err"=model error, "std"=standard deviation of measurement.
Figure 9.12: **Modeling fitting result of Hox C13.** Top left and right are expression levels of Oct4 and Nanog proteins for WT and KO context respectively, where blue line is Oct4 protein, green line is Nanog protein. Bottom left and right are expression levels of mRNA Hox C13 for WT and KO respectively. Blue line is the observed expression level of Hox C13, red dash line is the model prediction of Hox C13. "err"=model error, "std"=standard deviation of measurement.
Figure 9.13: **Modeling fitting result of Hox D3.** Top left and right are expression levels of Oct4 and Nanog proteins for WT and KO context respectively, where blue line is Oct4 protein, green line is Nanog protein. Bottom left and right are expression levels of mRNA Hox D3 for WT and KO respectively. Blue line is the observed expression level of Hox D3, red dash line is the model prediction of Hox D3. "err"=model error, "std"=standard deviation of measurement.
Figure 9.14: **Modeling fitting result of Hox D8.** Top left and right are expression levels of Oct4 and Nanog proteins for WT and KO context respectively, where blue line is Oct4 protein, green line is Nanog protein. Bottom left and right are expression levels of mRNA Hox D8 for WT and KO respectively. Blue line is the observed expression level of Hox D8, red dash line is the model prediction of Hox D8. "err"=model error, "std"=standard deviation of measurement.
Figure 9.15: **Modeling fitting result of Hox D9.** Top left and right are expression levels of Oct4 and Nanog proteins for WT and KO context respectively, where blue line is Oct4 protein, green line is Nanog protein. Bottom left and right are expression levels of mRNA Hox D9 for WT and KO respectively. Blue line is the observed expression level of Hox D9, red dash line is the model prediction of Hox D9. "err"=model error, "std"=standard deviation of measurement.
Figure 9.16: **Modeling fitting result of Hox C10 of motif type A.** Top left and right are expression levels of miRNA mmu-mir-33 for WT and KO context respectively. Middle left and right are expression levels of Oct4 and Nanog proteins for WT and KO context respectively, where blue line is Oct4 protein, green line is Nanog protein. Bottom left and right are expression levels of mRNA Hox C10 for WT and KO respectively. Blue line is the observed expression level of Hox C10, red dash line is the model prediction of Hox C10. "err"=model error, "std"=standard deviation of measurement.
Figure 9.17: **Modeling fitting result of Hox C10 of motif type A.** Top left and right are expression levels of miRNA mmu-mir-155 for WT and KO context respectively. Middle left and right are expression levels of Oct4 and Nanog proteins for WT and KO context respectively, where blue line is Oct4 protein, green line is Nanog protein. Bottom left and right are expression levels of mRNA Hox C10 for WT and KO respectively. Blue line is the observed expression level of Hox C10, red dash line is the model prediction of Hox C10. "err"=model error, "std"=standard deviation of measurement.
Figure 9.18: **Modeling fitting result of Hox C10 of motif type A.** Top left and right are expression levels of miRNA mmu-mir-143 for WT and KO context respectively. Middle left and right are expression levels of Oct4 and Nanog proteins for WT and KO context respectively, where blue line is Oct4 protein, green line is Nanog protein. Bottom left and right are expression levels of mRNA Hox C10 for WT and KO respectively. Blue line is the observed expression level of Hox C10, red dash line is the model prediction of Hox C10. "err"=model error, "std"=standard deviation of measurement.
Figure 9.19: **Modeling fitting result of Hox C10 of motif type A.** Top left and right are expression levels of miRNA mmu-mir-345 for WT and KO context respectively. Middle left and right are expression levels of Oct4 and Nanog proteins for WT and KO context respectively, where blue line is Oct4 protein, green line is Nanog protein. Bottom left and right are expression levels of mRNA Hox C10 for WT and KO respectively. Blue line is the observed expression level of Hox C10, red dash line is the model prediction of Hox C10. "err"=model error, "std"=standard deviation of measurement.
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Table 9.4: Motif B: Number of Validated miRNAs

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<th>protein G target</th>
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Table 9.5: Motif B: Validated MiRNAs Repressing Oct4

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<th>protein</th>
<th>class</th>
<th>miRNAs</th>
<th>protein</th>
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Table 9.6: Motif B: Validated Repressing GCNF

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Chapter 10

Condensation of Data

10.1 Distance of Expression Level of Two Molecules

Define an affine invariant distance for two vectors \( g_1 \) and \( g_2 \), denoted as \( D(g_1, g_2) \):

\[
D(g_1, g_2) = \sqrt{2(1 - corr(g_1, g_2))}
\]

where \( corr(g_1, g_2) \) is the correlation between \( g_1 \) and \( g_2 \).

Let \( \overline{g_1} = mean(g_1(t)), \overline{g_2} = mean(g_2(t)), p= \) number of time points,

\[
corr(g_1, g_2) = \frac{\frac{1}{p} \sum (g_1(t) - \overline{g_1})(g_2(t) - \overline{g_2})}{\sqrt{\frac{1}{p} \sum (g_1(t) - \overline{g_1})^2 \frac{1}{p} \sum (g_2(t) - \overline{g_2})^2}}
\]

So if we normalize the concentration \( g_1(t) \) to get normalized concentration \( U_n(t) \) as follows:

\[
U_n(t) = \frac{g_1(t) - \overline{g_1}}{\sum (g_1(t) - \overline{g_1})^2}
\]

Let \( U_n = (\frac{(g_1(t) - \overline{g_1})}{\sum (g_1(t) - \overline{g_1})^2})_t, U_m = (\frac{(g_2(t) - \overline{g_2})}{\sum (g_2(t) - \overline{g_2})^2})_t \). Then \( U_n \) and \( U_m \) are two unit vectors and are the normalized concentration vector for \( g_1 \) and \( g_2 \), the equation above can be rewritten.
as:

\[ \text{corr}(g_1, g_2) = U_n \cdot U_m \]

Let \( \theta \) be the angle between vector \( U_n \) and \( U_m \), then

\[ \cos(\theta) = \frac{U_n \cdot U_m}{\|U_n\| \|U_m\|} = U_n \cdot U_m \]

because \( U_n \) and \( U_m \) are unit vectors.

So \( D(g_1, g_2) = \sqrt{2(1 - \text{corr}(g_1, g_2))} = \sqrt{2(1 - \cos(\theta))} \), and since

\[ \cos(\theta) \approx 1 - \frac{\theta^2}{2} \]

as \( \theta \) is small.

Then

\[ D(g_1, g_2) \approx \sqrt{2(1 - (1 - \frac{\theta^2}{2}))} = \sqrt{\theta^2} = \theta \]

So if the angle of the two normalized trajectories is very small, then we will get their affine invariant distance small, too.

### 10.2 A Second Definition of Distance Between Two Vectors

Define a second affine invariant distance for two vectors \( g_1 \) and \( g_2 \), denoted as \( D(g_1, g_2) \):

\[ \tilde{D}(g_1, g_2) = \max(\left| \frac{g_1}{\max g_1} - \frac{g_2}{\max g_2} \right|) \]

If \( \tilde{D}(g_1, g_2) \) is sufficiently small, then \( g_1 \approx \alpha g_2 \). Since the equation for motif A will remains invariant by scale transformation, the upstream factors that are validated for mRNA \( G_1 \) can also be validated for mRNA \( G_2 \). Similarly, if \( \tilde{D}(m_1, m_2) \) is small enough, and \( m_1 \) is a validated miRNA that is predicted to degrade mRNA \( G \) by modeling motif B, then \( m_2 \) can also be validated to degrade mRNA \( G \).
10.3 Minimal Net Method to Condense the Data of miRNA

Given $N$ vectors of concentration in $\mathbb{R}^p$, where $p = 19$, we denote the vectors as $m_1, \ldots, m_N$, where $i$ is a positive integer recording the indices of the vectors, $i = 1, \ldots, N$. In general case, we simply denote the first vector in our list to be 1st, the 2nd vector in our list to be 2, and so fourth.

Let vector $A = [1, \ldots, N]$, set $AA = 1, \ldots, N$. Fix a small number $L$, which is considered to be the radius of a group of vectors in affine invariant distance.

The algorithm is as follows:

1. Let $DD_{j,k} = D(m_j, m_k), j, k = 1, \ldots, N$, find out the maximum value of the matrix $M$, which is $DD_{JK} = \max_{j,k}(DD_{jk})$. Denote a set $NET_1$, $NET_1 = \{J, K\}$; $NET_1$ is a set to store the indices of the miRNA concentration vectors, each of which we select is the center of each group. So now we first get two centers for two groups respectively. If $DD_{JK} < L$, denote a vector $n = [J, K]$, which is the vector form of set $NET_1$.

   Stop if $DD_{J,K} > L$, otherwise go to step 2.

2. In the $m^{th}$ iteration, suppose we have found out the set $NET_m$, $NET_m$ is a set to store the indices of the miRNA concentration vectors, each of which we select is the center of each group. Let’s call $NET_m$ ”center set” in short. Renew the vector $n=$vector form of set $NET_m$.

   Let

   $$S = AA \backslash NET_m = \{i| i \ is in \ AA \ but \ not \ in \ NET_m\}$$

   Renew $A=$vector form of $S$, then $A$ contains the indices of miRNA concentration vectors that have not been selected as centers.
Renew matrix $DD$, let $DD(j, k) = D(m_n_j, A_k)$, where $n_j$ is the $j^{th}$ element of vector $n$, $A_k$ is the $k^{th}$ element of vector $A$. So basically the matrix $DD$ now contains the distance between each concentration vector inside the "center" set and each concentration vector outside the "center" set.

Let $aa$, $bb$ be number indices satisfying:

$$DD_{aa,bb} = \max_k (\min_j (DD_{j,k}))$$

If $DD_{aa,bb} < L$, Then $NET_{m+1} = NET_m, bb$. $n = [Net(m + 1)]$. Stop if $DD_{aa,bb} > L$, otherwise continue and repeat step 2.

If we stop at $mm^{th}$ iteration, then the $1 \times (mm + 1)$ vector $n$ contains the $mm + 1$ centers for the $mm + 1$ groups. Denote these centers as $C_1, \ldots, C_{mm+1}$. We say that the miRNA $i$ is in group $m$ if $D(m_i, m_{C_m}) = \min_j (D(m_i, m_{C_j}))$. Thus we can get $mm + 1$ groups. We obtained 121 and 113 groups for the WT and KO data, in which 59 and 69 groups have more than 1 cardinals for WT and KO respectively. The diameter of miRNA cluster is no more than .15, the diameter of mRNA cluster is no more than 4% with respect to range 1.

The following are the names of the cluster presented on figure [10.1].
Figure 10.1: Normalized curves of a miRNA cluster
mmu-miR-293
mmu-miR-1
mmu-miR-101a
mmu-miR-101b
mmu-miR-150
mmu-miR-182
mmu-miR-183
mmu-miR-23a
mmu-miR-290
mmu-miR-291b-5p
mmu-miR-292-3p
mmu-miR-292-5p
mmu-miR-293
mmu-miR-29b
mmu-miR-92

10.4 Condensation for Data of mRNA

Theoretically we can use the Minimal Net Method to condense the mRNA data. However, the mRNA microarray data contains over 45000 records of observations, which will generate huge matrix in Minimal Net method that is not feasible for efficient computation.

We consider another simpler method specifically applicable for condensation of 3-time-point data of mRNAs. First we normalize the concentration of mRNA $G_n$, for $n = 1, \ldots, 45101$ as follows:

$$U_n(t) = \frac{g_n(t) - \bar{g}_n}{\sum (g_n(t) - \bar{g}_n)^2}$$
Since each normalized data is only one dimensional, we choose a unit vector \( e = [1, 0, 0] \).

Compute the dot product of \( e \) and \( U_j \), \( j = 1, \ldots, 45101 \),

\[
e \cdot U_j = ||e|| ||U_j|| \cos(\theta_j) = \cos(\theta_j)
\]

Where \( \theta_j \) is the angle between \( e \) and \( U_j \).

We can classify the 45101 observations into 100 groups first using the \( \cos \) values computed above, by calculating the quantiles for vector \( v = [\cos(\theta_1), \ldots, \cos(\theta_{45101})] \). Let \( Q(i/100) \) be the quantile for \( v \) at \( i/100 \), i.e. fix \( i \), the frequency that \( \cos(\theta_j) \leq Q(i/100) \) is \( i/100 \), \( i = 0, \ldots, 99 \), \( j = 1, \ldots, 45101 \).

Thus we obtained the 100 intervals \( [Q(i/100), Q((i + 1)/100)] \). Note that \( \arccos(Q(i/100)) \) can give us 2 distinct angles in \([0, 2\pi]\). Therefore, the mRNAs falling in the same specific subinterval by the \( \cos \) value, have to be divided into 2 groups. For a specific group \( i \), all the mRNAs \( j \) in this group satisfying:

\[
e \cdot U_j \in [Q(i/100), Q((i + 1)/100)], i = 0, \ldots, 99
\]

Fix mRNA \( G_j \) in group \( i \), and randomly select mRNA \( G_k \) in group \( i \). If

\[
corr(U_j - e, U_k - e) \geq 0
\]

Then together with the close \( \cos \) values, the trajectories of \( U_j \) and \( U_k \) are also very close, then we can put mRNA \( k \) and mRNA \( j \) together in the same group, name it \( G_i \). Otherwise,

\[
corr(U_j - e, U_k - e) < 0
\]

Then we put mRNA \( k \) to be in another group, name it \( G_{200-i} \).

In all there are 200 groups.

The above algorithm for condensation is especially for 3 dimensional mRNA data normalized by linear transformation. There is a more general way for grouping the huge mRNA
data efficiently, which is also applicable for higher dimension observations and scale normalization or other distance definition. We can first apply K-means clustering to get $K$ subsets of mRNA data with much smaller size, then use Minimal Net method for each subsets and refine the grouping.

10.5 Condensed ODE System For Concentration Profiles

As we have discussed in section 7.6, CKE models of both motif type A and B keep invariant under scale changes. With the condensation data under pairwise distance defined in a scale-relation way, the number of downstream factors can be greatly reduced, thus modeling global genetic network is feasible with much less computational cost.
Chapter 11

Conclusion and Further Discussion

In the biological context of Embryonic Stem Cells differentiation, we have separately modelled the 2 distinct modalities of the repressive functions of micro-RNAs in the post-transcriptional processes linked to differentiation regulators. This was achieved by first defining the formal structure of two types of interaction architectures (motifs A and B) linking micro-RNAs to subgroups of the mRNA genes these micro-RNAs are expected to target, and to the proteins associated to these mRNA genes. Then we have generated two quite long lists (called List$_A$ and List$_B$) of more than 10,000 concrete biologically plausible instances of potential motifs A and motifs B.

The plausibility of each one of these motif A or motif B instances was then evaluated by computerized parametric modeling, based on our two sets of microarray data and on adequate formal chemical kinetics equations (CKEs). We have sketched the formal derivation of 2 specific CKEs modelling by dynamic ODEs the interactions between the concentrations of different species of molecules involved in each motif.
This led to an architecture validation algorithm based on the quantified quality of fit between our optimally parametrized models and the corresponding microarray data.

To significantly narrow the List$_A$, we have naturally constrained the potentially interacting pairs of genes (M= miRNA, G=mRNA) involved in our selected motifs instances to be such that the potential targeting of gene G by gene M is actually predicted by one of the two biological reference tables miRanda and TargetScan.

For List$_B$, more than one miRNAs ($M_1, M_2, \ldots$) could possibly target a specific mRNA gene $G$ in order to inhibit the translation of gene $G$. In order to force the ratio of the number of parameters to the number of data points to remain as small as possible, we have deliberately restricted the motifs of List$_B$ to involve at most 3 microRNA repressors $M_1, M_2, M_3$, and to be potential targeting gene $G$ according to the tables miRanda or TargetScan.

For each motif in List$_A$ or List$_B$, the associated CKE model is then automatically parametrized in order to best fit our 2 sets of microarray data recording more than 20,000 genes expressions during ES-cells differentiation. One experiment involved ES-cells of wild type WT and the other experiment involved "GCNF-knockout" ES-cells. This computerized parameter estimation is implemented by our innovative fast algorithm, to handle in reasonable time the several thousands of motifs to be evaluated. On a laptop PC, our implementation of parameter estimation for a typical 9-parameters CKE model requires about 5 minutes of computing time. The estimation algorithm we have implemented also provides relatively high-quality optimization for the fit between model and data, by integrating both global and local cost minimization techniques, in contexts where plausible ranges of values for most of the unknown parameters are not available in the literature. This technique seems quite applicable to similar problems of parameter estimation in chemical kinetics modeling of molecular interactions.
After this optimized parametrization, the CKE models of each motif in List\textsubscript{A} or List\textsubscript{B} are tested by comparing the genes expression levels actually recorded in the WT and KO microarray data with the predicted numerical expression levels generated by the parametrized models.

We have set up adequate numerical tests to evaluate the sensitivity of model prediction to small changes of parameter values, and to analyze the sensitivity of parameter estimation to the measurements errors corrupting the microarray data sets. This approach is efficient but requires intensive computing time, so that the massive analysis of sensitivity results will be completed in future work. Our automated modeling and validation procedure of potential architectural motifs involving the potential repressive actions of micro-RNAs on key mRNA genes controlling the regulation of ES-cells differentiation has enabled us to generate much smaller final lists of validated motifs A and B. Our methodology and the associated intensive computations we have performed thus generate several interesting "motif A"-families of "model validated" interacting miRNA-mRNA pairs, as well as "motif B"-families of model validated groups of miRNAs inhibiting specific key proteins linked to differentiation regulation. All these model validated pairs of miRNA-mRNA genes are listed in the present work. These lists of validated miRNA-mRNA pairs or miRNAs-protein combinations should be of interest to efficiently circumscribe further biological experiments, by focusing gene expression recordings on much smaller sets of miRNAs than those predicted by the wide range biological reference tables miRanda or TargetScan. Moreover for any given mRNA gene G of specific interest, our results and methods should help narrow considerably the biological experiments to be performed in order to determine which miRNAs directly degrade gene G, and which miRNAs repress the translation of G. .

A key facilitating point to implement our modelization techniques is the availability of
expression levels for the proteins associated to the main mRNA genes of interest. These proteins expression levels are difficult to measure simultaneously for large sets of proteins. But we can still use our CKE models to determine whether the actual proteins whose expression levels are effectively measurable, are indeed transcriptional factors for a specific mRNA gene $G$.

Modeling very large microarray datasets is computationally quite expensive. We have hence sketched clustering methods to condense the 20,000 recorded expression levels profiles. This approach has of course been attempted before our work, but the main point is that we have carefully studied the mathematical compatibility of our CKE models with condensation of the profiles data. Since we have proved that the abstract form of our models is invariant by arbitrary multiple affine transformations of the profile data, we have made sure to constrain the distance of two expression levels profiles to be invariant by these types of affine transformations.

We have implemented a Minimal Net Clustering algorithm based on this distance. The number of CKEs to parametrize can be strongly reduced after condensation of the 20,000 profiles, and the affine invariance of our CKEs show that the condensed genes network can then still be modelled by similar CKEs. Even after condensation the exhaustive lists of potential motifs A and motifs B architectures still involve quite high numbers of architectures, due to combinatorial complexity. Hence rather than implementing blind modelling and parametrization, we have privileged the systematic focus on architectures involving shorter known lists of key regulatory mRNA genes linked to ES-cells differentiation, in close relation with the previous biological results and interesting hypotheses published in [2], where our microarray data sets on ES-cells had previously been analyzed by much more classical techniques.
Appendix: Table of Validated List of Motif B
Table 11.1: **List of miRNAs repressing Oct4 for Motif B**

<table>
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