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Reverse Engineering in Systems Biology: Towards Inference of a Gene Regulatory Network

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1 Introduction

With the ad ance of technolog) in the field of biolog), large scale sequencing data of genomes, or massi e gene e pression data of mRNA are a ailable, recently. High-throughput technology could thus measure plent, data. In particular, such as DNA microarra, technology enables us to handle massi e gene e pression data simultaneously during a single e periment. Here arises a question: ho should useful and information be mined from the data? For e ample, can e e tract or infer a gene regulator, net ork from the data obtained by biological e periment? So far, in order to ans er the question, some attempts ere carried out, let efficient method ha e not proposed.

In this paper, e introduce our ork, hich is an attempt to de elop an ad anced scheme for inferring a gene regulator) net ork, and for design of biological e periment, from such arious amounts of e perimental data. Although e ha e not let established quite useful and efficient scheme of inferring a gene net ork, this is an initial report to sho our acti ities up to this time.

2 Concept

Our concept for de eloping the scheme is to infer the most plausible and reliable gene net ork, pro iding a po erful and efficient re erse engineering method. Besides, from the ie point of biological most efficient all for utili ing the re erse engineering method is to predict or to erifly a putati e path all These results ill lead to design biological e periments, in hich order the should be carried out. In order to achie e this, e deal ith to profiles: one is time series e pression profiles, and the other is steadly state e pression profiles. Bly using both different the of data, e can e tract sophisticated and reliable net ork, leading to de elop a hybrid inferring slystem as briefly described in Figure 1. Needless to sall, e en either the data is lacked, e should be able to e tract consistent gene net ork. We are currently orking on de eloping se eral methods from different perspecti es, as are briefly introduced in ne t section.

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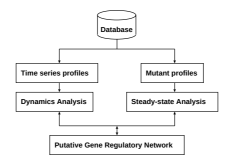


Figure 1: Schematic Representation of the Hibrid Inferring Sistem

3 Past Works

3.1 In silico Sampling and Screening

In pre ious paper [1], e ha e proposed a method, in silico Sampling and Screening, hich uses ild the and mutant e pression profiles to resol e plausible net ork candidates. The method is mainly consists to stage: sampling stage, and screening stage. First, in sampling stage, ild the profile is used so that candidate net ork topologies and their parameters are optimiled. Net, in screening stage, using mutant e pression profiles, final plausible candidate net orks are resol ed. Using 3-node net ork, 8 net ork topologies are e tracted as final candidate net orks.

3.2 Hypothesis Generator

The h) pothesis generator [3] is not only for identifying gene regulator) net orks from gene e pression profiles such as gene disruption and gene o er-e pression, but also for describing the e periment design and plans. Assuming the se eral e pression patterns of genes in *Drosophila* embryo, consistent net ork could be e tracted.

3.3 Application for Actual Expression Profiles

While the to methods mentioned abo e still rely on artificial fake data as the target, e has e also applied a method to infer a gene net ork for the actual e perimental data of Saccharomyces cerevisiae [2]. We employed clustering technique and neural net ork based model, and successfully achie ed to reduce the dimensionality and to infer net orks.

4 Concluding Remarks

We particularly focus on 1) data translation technique, and 2) parameter optimi ation technique currently, hich e think are both the most significant issues in establishing a sophisticated re erse engineering method. The technique of translating e pression data from ra e perimental data, hich makes easy to embed as input data in computer, ould accelerate to re eal underlying gene net ork. For fitting model—ith actual data, parameter optimi ation should be necessarly to get more reliable net ork. We belie e that these to techniques—ould allo—us to pro—ide an efficient re—erse engineering method.

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