Genome-Scale Gene Expression Profiles Mapped onto the Pathway and Genome Maps in KEGG

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1 Introduction

The emerging technology of DNA chips and microarrays makes it possible to simultaneously analyze the expression of many genes, such as the whole set of genes in the completely sequenced genome. We have been developing a system for network oriented analysis and visualization of such genome-scale gene expression data. It projects the time-series data of gene expression profiles on the functional relations map (KEGG pathway map) and the positional neighbors map (KEGG genome map), and uncovers underlying events in differential gene expression patterns between the reference and the perturbed state (e.g., the wild-type and a disruptant, or the control and an environmental shift)

2 Data and Methods

We have developed interfaces to the tools to adapt to any gene expression profiles data. In this report we use the numerical data for the microarray gene expression profiles of *Saccharomyces cerevisiae* [1] obtained by anonymous ftp [2]. The experiments involve two types of data, the environmental change of diauxic shift and the genetic change of deletion or overexpression of a transcription factor.

We use a part of KEGG (Kyoto Encyclopedia of Genes and Geneomes) [3] database; the positional neighbor data as represented by the GENOME graphical maps and the functional relation data as represented by the PATHWAY graphical maps. The basic colorization tool has been developed and it is made available in KEGG.

URL http://www.genome.ad.jp/kegg-bin/mk_point_pathway_multi_html for pathway maps URL http://www.genome.ad.jp/kegg-bin/mk_point_genome_multi_html for genome maps

The coloring strategy is as follows. The qualitative information in the data is represented by the difference in color, and the quantitative information is represented by the color depth. The change of two gene expression profiles between the reference state and the perturbed state is represented by two alternative colors for increased and decreased expression, a third color for no change, and a fourth color for no expression in both states. The time-series data representation is investigated such as GIF animation.

3 Results and Discussions

The publicly released microarray data of *Saccharomyces cerevisiae* was investigated. It contained a metabolic shift with genetic reprogramming, referred to as the diauxic shift from fermentation to respiration. This genetic reprogramming involved in basic cellular processes such as carbohydrate metabolism, nucleic acid and protein synthesis and energy metabolism. Fig. 1 shows the diauxic

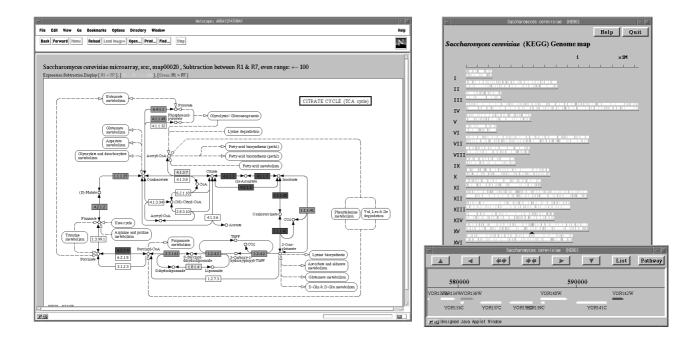


Figure 1: Gene expression profile mapped onto the PATHWAY (left) map and GENOME (right) map.

change that is automatically mapped on the pathway map of TCA cycle in KEGG. It is interesting to note that although there is no operon structure for the TCA cycle genes in *Saccharomyces cerevisiae* the bacterial genomes contain two separate operons, whenever they do have operons, that correspond to the upper and lower portions of the cycle as represented by different colors in Fig. 1. This suggests that there is a mechanism to cooperatively regulate the expression of genes that are far apart in the genome.

In general the correlation of gene expression patterns suggests functional relations and would assist to assign functions to hypothetical genes. By analyzing the positional correlation of synchronized expression of genes, common upstream activating sequences may also be found.

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