

Microarray Data Mining

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INTRODUCTION

Based on the concept of simultaneously studying the expression of a large number of genes, a DNA microarray is a chip on which numerous probes are placed for hybridization with a tissue sample. Biological complexity encoded by a deluge of microarray data is being translated into all sorts of computational, statistical, or mathematical problems bearing on biological issues ranging from genetic control to signal transduction to metabolism. Microarray data mining is aimed to identify biologically significant genes and find patterns that reveal molecular network dynamics for reconstruction of genetic regulatory networks and pertinent metabolic pathways.

BACKGROUND

The idea of microarray-based assays seemed to emerge as early as of the 1980s (Ekins & Chu, 1999). In that period, a computer-based scanning and image-processing system was developed to quantify the expression level in tissue samples of each cloned complementary DNA sequences spotted in a two-dimensional array on strips of nitrocellulose, which could be the first prototype of the DNA microarray. The microarray-based gene expression technology was actively pursued in the mid-1990s (Schena, Heller, & Thériault, 1998) and has seen rapid growth since then.

Microarray technology has catalyzed the development of the field known as functional genomics by offering high-throughput analysis of the functions of genes on a genomic scale (Schena et al., 1998). There are many important applications of this technology, including elucidation of the genetic basis for health and disease, discovery of biomarkers of therapeutic response, identification and validation of new molecular targets and modes of action, and so on. The accomplishment of decoding human genome sequence together with recent advances in the biochip technology has ushered in genomics-based medical therapeutics, diagnostics, and prognostics.

MAIN THRUST

The laboratory information management system (LIMS) keeps track of and manages data produced from each step in a microarray experiment, such as hybridization, scanning, and image processing. As microarray experiments generate a vast amount of data, the efficient storage and use of the data require a database management system. Although some databases are designed to be data archives only, other databases such as ArrayDB (Ermolaeva, Rastogi, & Pruitt, 1998) and Argus (Comander, Weber, Gimbrone, & Garcia-Cardena, 2001) allow information storage, query, and retrieval, as well as data processing, analysis, and visualization. These databases also provide a means to link microarray data to other bioinformatics databases (e.g., NCBI Entrez systems, Unigene, KEGG, and OMIM). The integration with external information is instrumental to the interpretation of patterns recognized in the gene-expression data. To facilitate the development of microarray databases and analysis tools, there is a need to establish a standard for recording and reporting microarray gene expression data. The MIAME (Minimum Information about Microarray Experiments) standard includes a description of experimental design, array design, samples, hybridization, measurements, and normalization controls (Brazma, Hingamp, & Quackenbush, 2001).

Data Mining Objectives

Data mining addresses the question of how to discover a gold mine from historical or experimental data, particularly in a large database. The goal of data mining and knowledge discovery algorithms is to extract implicit and previously unknown nontrivial patterns, regularities, or knowledge from large data sets that can be used to improve strategic planning and decision making. The discovered knowledge capturing the relations among the variables of interest can be formulated as a function for making prediction and classification or as a model for understanding the problem in a given domain. In the context of microarray data, the objectives are identifying significant genes and finding gene expression pat-

terns associated with known or unknown categories. Microarray data mining is an important topic in bioinformatics, dealing with information processing on biological data, particularly genomic data.

Practical Factors Prior to Data Mining

Some practical factors should be taken into account prior to microarray data mining. First of all, microarray data produced by different platforms vary in their formats and may need to be processed differently. For example, one type of microarray with cDNA as probes produces ratio data from two channel outputs, whereas another type of microarray using oligonucleotide probes generates nonratio data from a single channel. Not only may different platforms pick up gene expression activity with different levels of sensitivity and specificity, but also different data processing techniques may be required for different data formats.

Normalizing data to allow direct array-to-array comparison is a critical issue in array data analysis, because several variables in microarray experiments can affect measured mRNA levels (Schadt, Li, Ellis, & Wong, 2001; Yang, Dudoit, & Luu, 2002). Variations may occur during sample handling, slide preparation, hybridization, or image analysis. Normalization is essential for correct microarray data interpretation. In simple ways, data can be normalized by dividing or subtracting expression values by a representative value (e.g., mean or median in an array) or by taking a linear transformation to zero mean and unit variance. As an example, data normalization in the case of cDNA arrays may proceed as follows: The local background intensity is subtracted from the value of each spot on the array; the two channels are normalized against the median values on that array; and the Cy5/Cy3 fluorescence ratios and \log_{10} -transformed ratios are calculated from the normalized values. In addition, genes that do not change significantly can be removed through a filter in a process called *data filtration*.

Differential Gene Expression

To identify genes differentially expressed across two conditions is one of the most important issues in microarray data mining. In cancer research, for example, we wish to understand what genes are abnormally expressed in a certain type of cancer, so we conduct a microarray experiment and collect the gene expression profiles of normal and cancer tissues, respectively, as the control and test samples. The information regarding differential expression is derived from comparing the test against the control sample.

To determine which genes are differentially expressed, a common approach is based on fold-change; in this approach, we simply decide a fold-change threshold (e.g., 2C) and select genes associated with changes greater than that threshold. If a cDNA microarray is used, the ratio of the test over control expression in a single array can be converted easily to fold change in both cases of up-regulation (induction) and down-regulation (suppression). For oligonucleotide chips, fold-change is computed from two arrays, one for test and the other for control sample. In this case, if multiple samples in each condition are available, the statistical *t*-test or Wilcoxon tests can be applied, but the catch is that the Bonferroni adjustment to the level of significance on hypothesis testing would be necessary to account for the presence of multiple genes. The *t*-test determines the difference in mean expression values between two conditions and identifies genes with significant difference. The nonparametric Wilcoxon test is a good alternative in the case of non-Gaussian data distribution. SAM (Significance Analysis of Microarrays) (Tusher, Tibshirani, & Chu, 2001) is a state-of-the-art technique based on balanced perturbation of repeated measurements and minimization of the false discovery rate.

Coordinated Gene Expression

Identifying genes that are co-expressed across multiple conditions is an issue with significant implications in microarray data mining. For example, given gene expression profiles measured over time, we are interested in knowing what genes are functionally related. The answer to this question also leads us to deduce the functions of unknown genes from their correlation with genes of known functions. Equally important is the problem of organizing samples based on their gene expression profiles so that distinct phenotypes or disease processes may be recognized or discovered.

The solutions to both problems are based on so-called cluster analysis, which is meant to group objects into clusters according to their similarity. For example, genes are clustered by their expression values across multiple conditions; samples are clustered by their expression values across genes. The issue is the question of how to measure the similarity between objects. Two popular measures are the Euclidean distance and Pearson's correlation coefficient. Clustering algorithms can be divided into hierarchical and nonhierarchical (partitional). Hierarchical clustering is either agglomerative (starting with singletons and progressively merging) or divisive (starting with a single cluster and progressively breaking). Hierarchical agglomerative clustering is most commonly used in the

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